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(57) Abstract

Novel hybridisation assay probes and mixtures of such probes for detecting a target sequence of one or more mycobacteria optionally present in a sample. The probes may suitably be directed to target sequences of mycobacterial rDNA, precursor rRNA, or rRNA, said probes being capable of forming detectable hybrids. The probes are in particular directed to mycobacterial rDNA, to precursor rRNA, or to 23S, 16S or 5S rRNA. The probes are useful for detecting the organisms in test samples such as sputum, laryngeal swabs, gastric lavage, bronchial washings, biopsies, aspirates, expectorates, body fluids (spinal, pleural, pericardial, synovial, blood, pus, bone marrow), urine, tissue sections as well as food samples, soil, air and water samples, and cultures thereof.

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NOVEL PROBES FOR THE DETECTION OF MYCOBACTERIA

The present invention relates to novel probes and to mixtures of such probes, in addition to the design, construction and use of such novel probes or mixtures thereof for detecting a target sequence of one or more mycobacteria, which probes are capable of detecting such organism(s) optionally present in a test sample, e.g. sputum, laryngeal swabs, gastric lavage, bronchial washings, biopsies, aspirates, expectorates, body fluids (spinal, pleural, pericardial, synovial, blood, pus, bone marrow), urine, tissue sections as well as food samples, soil, air and water samples and cultures thereof. The invention relates in particular to novel probes and mixtures thereof for detecting the presence of one or more mycobacteria of the Mycobacterium tuberculosis Complex (MTC) and for detecting the presence of one or more mycobacteria other than mycobacteria of the Mycobacterium tuberculosis Complex (MOTT). The invention further relates to diagnostic kits comprising one or more of such probes. The probes of the present invention are surprisingly able to penetrate the cell wall of the mycobacteria, thus making possible the development of fast an easy-performed in situ protocols.

BACKGROUND OF THE INVENTION

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Tuberculosis is a very life-threatening and highly epidemic disease which is caused by infection with Mycobacterium tuberculosis. Tuberculosis is presently the predominant infectious cause of morbidity and mortality world-wide, and is estimated to kill about three million people annually. WHO estimates that the annual number of new cases of tuberculosis will increase from 7.5 million in 1990 to 10.2 million in 2000, an escalation that will result in approximately 90 million new cases during this decade. It is furthermore estimated that 30 million people will die from tuberculosis during the 1990s, which equals one quarter of preventable deaths among adults.

The prevalence of tuberculosis has been very high in the poorer parts of the world such as Asia, Africa and South-America, but in recent years an increase has also been observed in industrialised countries. This appears to be due to an interaction of various factors including i.a. patterns of migration, poorly organised tuberculosis programmes and nutrition problems. Furthermore, a serious threat will arise from the emergence of new strains that are drug resistant or worse, multi-drug resistant.

Mycobacteria are often divided into tuberculous mycobacteria, i.e. mycobacteria of the Mycobacterium tuberculosis Complex (MTC), and non-tuberculous mycobacteria, i.e. mycobacteria other than those of the Mycobacterium tuberculosis Complex (MOTT). The MTC

group comprises apart from M. tuberculosis, M. bovis, M. africanum and M. microti. Mycobacteria of the MOTT group are not normally pathogenic to healthy individuals but may cause disease in immunocompromised individuals, e.g. individuals infected with HIV. Clinical relevant mycobacteria of the MOTT group are in particular M. avium, M. intracellulare, M. kansasii and M. gordonae, but also M. scrofulaceum, M. xenopi and M. fortuitum.

M. avium and M. intracellulare together with M. paratuberculosis and M. lepraemurium constitute the Mycobacterium avium Complex (MAC). Extended with M. scrofulaceum, the group is named Mycobacterium avium -intracellulare -scrofulaceum Complex (MAIS).

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It is well-known that treatment of mycobacterial infections with antibiotics may lead to the emergence of drug resistant strains. Many antibiotic drugs excert their effects by interfering with protein synthesis or with transcription. Studies of the molecular mechanisms underlying certain antibiotic resistance phenotypes in clinical mycobacterium isolates have revealed mutations in rRNA genes. The development of resistance because of mutation(s) located in the rRNA gene is likely to occur since slow-growing mycobacteria have only a single rRNA operon. All mycobacteria populations comprise a minority of drug resistant mutants that have arisen by spontaneous mutation. These mutated mycobacteria do normally not survive particularly well, but, when single-drug therapy is offered as treatment, the drug susceptible bacteria are killed, and only the resistant mutants will survive and multiply, and, thus at some point, constitute the majority of the mycobacterial population. The selection of drug resistant bacteria due to inadequate drug therapy leads to a state of so-called "acquired drugresistance". In contrast, "primary drug-resistance" is used to characterise a situation where drug-resistant mycobacteria can be isolated from a patient who has never been treated for mycobacterial infection, and has become infected with drug-resistant mycobacteria from an individual suffering from infection with an acquired drug resistant bacterium.

Today, drug-resistance is determined primarily phenotypically by culturing clinical samples, in which presence of mycobacteria have been demonstrated, in the presence of the individual drugs. This is unfortunately a very slow and time-consuming procedure as the result of the drug-resistance studies depends on the growth rate of the mycobacteria, which are well-known to be slow. Thus, the result is not available until after several weeks.

Although the incidence of drug-resistance is, at least not yet, very common, it is nevertheless very important that resistant strains are identified and eradicated. Therefore, it is of major importance to find a reliable and rapidly performed method of diagnosing drug-resistance.

Presently, the detection of mycobacteria by microscopy is the most prevalent method for

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diagnosis. The sample (e.g. an expectorate) is stained for the presence of acid-fast bacilli using e.g. Ziehl-Neelsen staining. However, staining for acid-fast bacilli does not provide the necessary information about the type of infection, only whether acid fast bacilli are present in the sample, and this is in itself not sufficient information for establishing a diagnosis. Samples positive for acid fast bacilli, may subsequently be cultured in order to be able to perform species identification.

Since Ziehl-Neelsen staining cannot be used to determine whether the infection is caused by mycobacteria of the MTC group or mycobacteria other than mycobacteria of the MTC group, a positive staining frequently leads to very costly isolation of all the patients with suspected M. tuberculosis infection as well as treatment with medicaments to which the patient may not even respond.

Since the sensitivity of acid fast staining is only approximately 10⁴-10⁵ per ml smear negative samples should also be cultured as culture-based tests are sensitive, and as it may be possible to detect 10-100 organisms per sample, but the result is not available before up to 8 weeks of culturing. Likewise, information about drug susceptibility is not available until after 1-3 weeks of further testing.

Different solid or liquid media (Loewenstein Jensen slants and Dubos broth) have traditionally been used for culturing of mycobacteria-containing samples. Newer media include ESP Myco Culture System (Difco), MB/BacT (Organon Teknika), BacTec (Becton Dickinson) and MGIT (Becton Dickinson). These test media are based on colourmetric or fluorometric detection of carbon dioxide or oxygen produced by mycobacterial metabolism, and adapted to automated systems for large scale testing.

Species identification is presently carried out following culturing using traditional biochemical methods or probe hybridisation assays (e.g. AccuProbe by Gen-Probe Inc., USA). There is, therefore, an increasing need for means allowing a more rapid distinction between mycobacteria of the MTC group and mycobacteria other than those of the MTC group, and for further species identification of those especially mycobacteria other than those of the MTC group.

A number of new attempts to replace the culture-based methods relies on molecular amplification technology. Several methods have emerged, among them the polymerase chain reaction (PCR), the ligase chain reaction and transcription mediated amplification. The basic principle of amplification methods is that a specific nucleic acid sequence of the mycobacteria is amplified to increase the copy number of the specific sequence to a level where the

amplicon may be detectable. In principle, the methods offers the possibility of detecting only one target sequence, thus, in principle, making detection of mycobacteria present at low levels possible. However, it has become clear that the target amplification methods cannot replace culture-based methods as only samples which are positive by staining for acid fast bacilli (AFB) give a satisfactory sensitivity. Furthermore, specific problems exist for each method. The PCR method may give false negative results due to the presence of inhibitors such as haemoglobin. Another problem arises from cross-contamination of negative specimens and/or reagents with amplified nucleic acid present in the laboratory environment leading to false positive results. A disadvantage is that costly reagents are needed for performing these tests. Furthermore, specialised instrumentation is required, making these tests mainly useful in large specialised laboratories, and generally not applicable in smaller clinical laboratories.

Nucleic acid probes for detecting rRNA of mycobacteria have been described in for example US 5 547 842, EP-A 0 572 120 and US 5 422 242.

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Considering the perspective and impact the disease has, the development of rapid and preferably easy-performed and further economic feasible diagnostic detection tests are of utmost importance and would be a very valuable tool in the fight against the spread of tuberculosis.

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Peptide nucleic acids are pseudo-peptides with DNA-binding capability. The compounds were first reported in the early nineties in connection with a series of attempts to design nucleotide analogues capable of hybridising, in a sequence-specific fashion, to DNA and RNA, cf. WO 92/20702.

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Hybridisation of peptide nucleic acid probes to DNA and to RNA has been shown to obey the Watson-Crick base pairing rules, and peptide nucleic acid probes have been found to hybridise to a DNA or a RNA target with higher affinity and specificity than the nucleic acid counterparts. These properties are ascribed to the uncharged, as opposed to the charged, structure of the peptide nucleic acid and DNA or RNA backbones, respectively, and to the high conformational flexibility of the peptide nucleic acid molecules. These features - together with the documented stability of peptide nucleic acid towards a variety of naturally occurring nucleases and proteases that usually degrade DNA, RNA or proteins - invite for use of peptide nucleic acid probes as antisense therapeutic agents and opens potentially important applications in diagnostics.

SUMMARY OF THE INVENTION

The present invention relates to novel peptide nucleic acid probes and to mixtures of such probes for detecting a target sequence of one or more mycobacteria optionally present in a sample. In accordance with claim 1, the probes are directed to target sequences of mycobacterial rRNA, genomic sequences corresponding to said rRNA (rDNA) and precursor rRNA. rRNA is present in a high number of copies in each cell, and is hence a well suited target. The probes are, as defined in claim 2, suitably directed to target sequences of mycobacterial rDNA, precursor rRNA, or 23S, 16S or 5S rRNA.

Thus, in a first aspect, the invention features a hybridisation assay probe and a mixture of such probes for detecting a target sequence of one or more mycobacteria in accordance with claim 1 and 2. Under appropriate stringency conditions, Such probes should not to any significant degree cross-react with ribosomal nucleic acid from other not relevant organisms, present in the test sample, in particular other mycobacteria. Cross-reactivity to organisms that are unlikely to be present in the sample may not be of importance. In in situ assays implying examination by microscopy, it is further possible to distinguish mycobacteria from other bacteria based on the morphology of these bacilli.

The invention also relates to peptide nucleic acid probes in accordance with claim 3 for obtaining a target sequence and in accordance with claim 4 for obtaining a probe.

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In another aspect, the invention relates to novel peptide nucleic acid probes for detecting a target sequence of one or more mycobacteria of the MTC group, and one or more mycobacteria other than mycobacteria of the MTC group, which probes comprise from 6 to 30 polymerised peptide nucleic acid moieties (claim 5). Suitable probes of formula (I) are claimed in claim 6.

Claims 7 to 10 and 15 to 24 relate to probes or mixtures of such probes for detecting a target sequence of one or more mycobacteria of the MTC group. Claims 11 to 13 and 15 to 24 relate to probes or mixtures of such probes for detecting a target sequence of one or more mycobacteria other than mycobacteria of the MTC group (MOTT group). Claim 14 relates specifically to probes for detecting drug resistant mycobacteria. Claims 25 to 27 relate to the use of such probes or mixtures thereof.

In accordance with claims 28 to 34, the present invention also relates to a method for detecting the presence of mycobacteria.

In yet another aspect, the present invention relates to a kit (claim 35 and 36) comprising at least one peptide nucleic acid probe as defined in claims 1 to 24.

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Mycobacteria are characterised by a complex cell wall which contains myolic acids, complex waxes and unique glycolipids. It is generally recognised by those skilled in the art that this wall provides mycobacteria with extreme resistance to chemical and physical stress as compared to other bacteria, and, accordingly, makes them very difficult to penetrate and lyse. The low permeability of the cell wall is considered to be the main reason for the fact that only very few drugs are effective in the treatment of tuberculosis and other mycobacterial infections. As described in US 5 582 985, the wall appears further to prevent penetration by nucleic acid probes. Even with short probes (shorter than 30 nucleic acids), specific staining is low or often non-existent. Protocols that allow DNA probes to be used for in situ hybridisation to mycobacterial species are described in US 5 582 985. However, these protocols require dewaxing of the mycobacterial cell wall with xylene and further enzymatic treatment prior to the hybridisation step in order to make the mycobacterial cell wall permeable to the DNA probes.

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The problems set forth above have surprisingly been completely solved by the use of peptide nucleic acid probes. It has, surprisingly, been found that the peptide nucleic acid probes are able to penetrate the cell wall of the mycobacteria, and furthermore that this is taking place rapidly. The person skilled in the art would arrive at the conviction that it would be necessary to heavily treat the mycobacteria before hybridisation is carried out. Thus, based on the available prior art, there is a strong prejudice against carrying out hybridisation without prior destruction of the mycobacterial cell wall. The inventors have shown that this is indeed and unexpectedly possible. It has been demonstrated that the probes of the present invention are able to hybridise to mycobacterial precursor rRNA and rRNA without harsh treatment of the mycobacterial cells, thus avoiding a risk of interfering with the morphology of the cells. Using the present probes, specific and easy detection and, subsequently, diagnosis of tuberculosis and other mycobacterial infections are thus possible.

BRIEF DESCRIPTION OF THE FIGURES

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Alignments of rDNA sequences of M. tuberculosis (as a representative of the MTC group) and important closely related species thereto, including M. avium (as a representative of the mycobacteria other than those of the MTC group) and important closely related species thereto for the 23S, 16S and/or 5S rRNA genes have been made (Figures 1A-1J, 2A-2D, 3, 4A-4L and 5A-B). The alignment for M. bovis and M. intracellulare are partly based on public available sequences and partly on sequences obtained by sequencing performed at DAKO A/S.

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Alignment for the MTC group (23S rDNA)

Figures 1A-1J show alignments of 23S rDNA sequences of M. tuberculosis (GenBank entry GB:MTCY130, accession number Z73902), M. avium (GenBank entry GB:MA23SRNA, accession number X74494), M. paratuberculosis (GenBank entry GB:MPARRNA, accession number X74495), M. phlei (GenBank entry GB:MP23SRNA, accession number X74493), M. leprae (GenBank entry GB:ML5S23S, accession number X56657), M. gastri (GenBank entry GB:MG23SRRNA, accession number Z17211), M. kansasii (GenBank entry GB:MK23SRRNA, accession number Z17212), and M. smegmatis (GB:MS16S23S5. accession number Y08453). Preferred peptide nucleic acid probes should enclose at least one nucleobase complementary to a nucleobase of M. tuberculosis 23S rRNA within positions 10 149-158, 220-221, 328-361, 453-455, 490-501, 637-660, 706-712, 762-789, 989, 1068-1072, 1148, 1311-1329, 1361-1364, 1418, 1563-1570, 1627-1638, 1675-1677, 1718, 1734-1740, 1967-1976, 2403-2420, 2457-2488, 2952-2956, 2966-2969, 3000-3003, and 3097-3106 of the alignment (indicated by heavy frames). Differences between the sequences of M. avium, M. phlei, M. leprae, M. paratuberculosis, M. gastri and M. kansasii and that of M. tuberculosis in 15 the alignment are indicated by light frames.

Alignment for the MTC group (16S rDNA)

Figures 2A-2D show alignments of 16S rDNA sequences of M. tuberculosis (GenBank entry 20 GB:MTU16SRN, accession number X52917), M. bovis (GenBank entry GB:MSGTGDA. accession number M20940), M. avium (GenBank entry GB:MSGRRDA, accession number M61673), M. intracellulare (GenBank entry GB:MIN16SRN, accession number X52927), M. paratuberculosis (GenBank entry GB:MSGRRDH, accession number M61680), M. scrofulaceum (GenBank entry GB:MSC16SRN, accession number X52924), M. leprae (GenBank entry GB:MLEP16S1, accession number X55587), M. kansasii (GenBank entry 25 GB:MKRRN16, accession number X15916), M. gastri (GenBank entry GB:MGA16SRN, accession number X52919), M. gordonae (GenBank entry GB:MSGRR16SI, accession number M29563) and M. marinum (GenBank entry GB:MMA16SRN, accession number X52920). Preferred peptide nucleic acid probes should enclose at least one nucleobase 30 complementary to a nucleobase of M. tuberculosis 16S rRNA within positions 76-79, 98-101, 135-136, 194-201, 222-229, 242, 474, 1136-1145, 1271-1272, 1287-1292, 1313, and 1334 of the alignment (indicated by heavy frames). Differences between the sequences of M. bovis, M. avium, M. intracellulare, M. paratuberculosis, M. scrofulaceum, M. leprae, M. kansasii, M. gastri, M. gordonae and M. marinum, and that of M. tuberculosis in the alignment are indicated 35 by light frames.

Alignment for the MTC group (5S rDNA)

Figure 3 shows alignments of 5S rDNA sequences of M. tuberculosis (GenBank entry

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GB:MTDNA16S, accession number x75601), M. bovis (GenBank entry GB:MBRRN5S, accession number X05526), M. phlei (GenBank entry GB:MP5SRRNA, accession number X55259), M. leprae (GenBank entry GB:ML5S23S, accession number X56657), and M. smegmatis (GenBank entry GB:MS16S23S5, accession number Y08453). Preferred peptide nucleic acid probes should enclose at least one nucleobase complementary to a nucleobase of M. tuberculosis 5S rRNA within positions 86-90 of the alignment (indicated by heavy frame). Differences between the sequences of M. bovis, M. phlei, M. leprae, M. smegmatis and M. luteus and that of M. tuberculosis in the alignment are indicated by light frames,

10 Alignment for Mycobacteria other than those of the MTC group (23S rDNA) Figures 4A-4L show alignments of 23S rDNA sequences of M. avium (GenBank entry GB:MA23SRNA, accession number X74494), M. paratuberculosis (GenBank entry GB:MPARRNA, accession number X74495), M. tuberculosis (GenBank entry GB:MTCY130, accession number Z73902), M. phlei (GenBank entry GB:MP23SRNA, accession number 15 X74493), M. leprae (GenBank entry GB:ML5S23S, accession number X56657), M. gastri (GenBank entry GB:MG23SRRNA, accession number Z17211), M. kansasii (GenBank entry GB:MK23SRRNA, accession number Z17212), and M. smegmatis (GB:MS16S23S5, accession number Y08453). Preferred peptide nucleic acid probes should enclose at least one nucleobase complementary to a nucleobase of M. avium 23S rRNA within positions 99-101. 20 183, 261-271, 281-284, 290-293, 327-335, 343-357, 400-405, 453-462, 587-599, 637-660, 704-712, 763-789, 1060-1074, 1177-1185, 1259-1265, 1311-1327, 1345-1348, 1361-1364, 1556-1570, 1608-1613, 1626-1638, 1651-1659, 1675-1677, 1734-1741, 1847-1853, 1967-1976, 2006-2010, 2025-2027, 2131-2232, 2252-2255, 2396-2405, 2416-2420, 2474-2478, 2687, 2719, 2809, 3062-3068, and 3097-3106 of the alignment (indicated by heavy frames). Differences between the sequences of M. paratuberculosis, M. tuberculosis, M. phlei, M. leprae, M. gastri, M. kansasii, and M. smegmatis and that of M. avium in the alignment are indicated by light frames.

Alignment for Mycobacteria other than those of the MTC group (16S rDNA)
 Figures 5A-5B show alignments of 16S rDNA sequences of M. avium (GenBank entry GB:MSGRRDA, accession number M61673), M. intracellulare (GenBank entry GB:MIN16SRN, accession number X52927), M. paratuberculosis (GenBank entry GB:MSGRRDH, accession number M61680), M. scrofulaceum (GenBank entry GB:MSC16SRN, accession number X52924), M. tuberculosis (GenBank entry GB:MTU16SRN, accession number X52917), M. bovis (GenBank entry GB:MSGTGDA, accession number M20940), M. leprae (GenBank entry GB:MLEP16S1, accession number X55587), M. kansasii (GenBank entry GB:MKRRN16, accession number X15916), and M. gastri (GenBank entry GB:MSGRR16SI, accession number X658RR16SI, accession number X658RR16SI,

accession number M29563), and M. marinum (GenBank entry GB:MMA16SRN, accession number X52920). Preferred peptide nucleic acid probes should enclose at least one nucleobase complementary to a nucleobase of M. avium 16S rRNA within positions 135-136, 472-475, 1136-1144, 1287-1292, 1313, and 1334 of the alignment (indicated by heavy frames). Differences between the sequences of M. intracellulare, M. paratuberculosis, M. scrofulaceum, M. tuberculosis, M. bovis, M. leprae, M. kansasii, and M. gastri and that of M. avium in the alignment are indicated by light frames.

Drug-resistance

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- Figure 6 shows a partial M. avium 23S rDNA sequence including positions 2550 to 2589 of GenBank entry X74494. Bases in positions where deviations from the wild-type sequence have been correlated with macrolide-resistance are framed. Positions 2568 and 2569 in the figure correspond to positions 2058 and 2059, respectively, of E. coli 23S rRNA.
- Figure 7 shows a partial M. tuberculosis 16S rDNA sequence including positions 441 to 491 and 843 to 883 of GenBank entry X52917. Bases in positions where deviations from the wild-type sequence have been correlated with resistance to streptomycin are framed. Positions 452, 473, 474, 477, 865, and 866 in the figure correspond to positions 501, 522, 523, 526, 912, and 913, respectively, of E.coli 16S rRNA.

SPECIFIC DESCRIPTION

The present invention provides novel probes for use in rapid and specific, sensitive hybridisation based assays for detecting a target sequence of one or more mycobacteria, which target sequence is located in the mycobacterial rDNA, precursor rRNA, or in the 23S, 16S or 5S rRNA. The probes to be used in accordance with the present invention are peptide nucleic acid probes. Peptide nucleic acids are non-naturally occurring polyamides or polythioamides which can bind to nucleic acids (DNA and RNA). Such compounds are described in e.g. WO 92/20702.

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We have identified suitable variable regions of the target nucleic acid by comparative analysis of generally available rDNA sequences and sequences obtained by sequencing as described above. Computers and computer programs, which have been used for the purposes disclosed herein, are commercially available. From such alignments, possibly suitable probes can be identified. The alignments are thus a useful guideline for designing probes with desired characteristics.

When designing the probes, due regard should be taken to the assay conditions under which

the probes are to be used. Stringency is chosen so as to maximise the difference in stability between the hybrid formed with the target nucleic acid and that formed with the non-target nucleic acid. It will typically be necessary to choose high stringency conditions for probes where the specificity depends on only one mismatch to non-target sequences. The more mismatches to non-target sequences, the less demand for high stringency conditions.

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Furthermore, probes should be designed so as to minimise the stability of probe-non-target nucleic acid hybrids. This may be accomplished by minimising the degree of complementarity to non-target nucleic acid, i.e. by designing the probe to span as many destabilising mismatches as possible, and/or to include as many additions/deletions relative to the target sequence as possible. Whether a probe is useful for detecting a particular mycobacterial species depends to some degree on the difference between the thermal stability of probetarget hybrids and probe:non-target hybrids. For rRNA targets, however, the secondary structure of the region of the rRNA molecule in which the target sequence is located may also be of importance. The secondary structure of a probe should also be taken into consideration. Probes should be designed so as to minimise their proclivity to form hairpins, self-dimers, and pair-dimers if a mixture of two or more probes is used.

Mismatching bases in hybrids formed between peptide nucleic acid probes and nucleic acids result in a higher thermal instability than mismatching bases in nucleic acid duplexes of the same sequences. Thus, the peptide nucleic acid probes exhibit a greater specificity for a given target nucleic acid sequence than a traditional nucleic acid probe, which is seen as a greater difference in T_m values for probe-target hybrids and probe-non-target hybrids. The sensitivity and specificity of a peptide nucleic acid probe will also depend on the hybridisation conditions used.

The primary concern regarding the length of the peptide nucleic acid probes is the warranted specificity, i.e. which length provides sufficient specificity for a particular application. The optimal length of a peptide nucleic acid probe comprising a particular site with differences in base composition, e.g. among selected regions of mycobacterial rRNA, is a compromise between the general pattern that longer probes ensure specificity and shorter probes ensure that the destabilising differences in base composition constitute a greater portion of the probe. Also, due regard must be paid to the conditions under which the probes are to be used.

Peptide nucleic acid sequences are written from the N-terminal end of the sequence towards the C-terminal end. A free (unsubstituted) N-terminal end or an N-terminal end terminating with an amino acid is indicated as H, and a free C-terminal end is indicated as NH₂ (a carboxamide group). Peptide nucleic acids are capable of hybridising to nucleic acid

sequences in two orientations, namely in antiparallel orientation and in parallel orientation. The peptide nucleic acid is said to hybridise in the antiparallel orientation when the N-terminal end of the peptide nucleic acid is facing the 3' end of the nucleic acid sequence, and to hybridise in the parallel orientation when the C-terminal end of the peptide nucleic acid is facing the 5' end of the nucleic acid sequence. In most applications, hybridisation in the antiparallel orientation is preferred as the hybridisation in the parallel orientation takes place rather slowly and as the formed duplexes are not as stable as the duplexes having antiparallel strands. Triplex formation with a stoichiometry of two peptide nucleic acid strands and one nucleic acid strand may occur if the peptide nucleic acid has a high pyrimidine content. Such triplexes are very stable, and probes capable of forming triplexes may thus be suitable for certain applications.

Mainly because the peptide nucleic acid strand is uncharged, a peptide nucleic acid-nucleic acid-duplex will have a higher T_m than the corresponding nucleic acid-nucleic acid-duplex. Typically there will be an increase in T_m of about 1 °C per base pair at 100 mM NaCl depending on the sequence (Egholm et al. (1993), Nature, 365, 566-568).

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In contrast to DNA-DNA-duplex formation, no salt is necessary to facilitate and stabilise the formation of a peptide nucleic acid-DNA or a peptide nucleic acid-RNA duplex. The T_m of the peptide nucleic acid-DNA-duplex changes only little with increasing ionic strength. Typically for a 15-mer, the T_m will drop only 5 °C when the salt concentration is raised from 10 mM NaCl to 1 M NaCl. At low ionic strength (e.g. 10 mM phosphate buffer with no salt added), hybridisation of a peptide nucleic acid to a target sequence is possible under conditions where no stable DNA-DNA-duplex formation occurs. Furthermore, target sites that normally are inaccessible can be made more readily accessible for hybridisation with peptide nucleic acid probes at low salt concentration as the secondary and tertiary structure of nucleic acids are destabilised under such conditions. Using peptide nucleic acid probes, a separate destabilising step or use of destabilising probes may not be necessary to perform,

rRNA is essential for proper function of the ribosomes and thus the synthesis of proteins. The genes encoding the rRNAs are in eubacteria located in an operon in which the small subunit RNA gene, the 16S rRNA gene, is located nearest the 5' end of the operon, the gene for the large subunit RNA, the 23S rRNA gene, is located distal to the 16S rRNA gene and the 5S rRNA gene is located nearest the 3' end of the operon. The three genes are separated by spacer regions in which tRNA genes may be found, however, there are none in M. tuberculosis. The primary transcript of the eubacterial rRNA operon is cleaved by RNaseIII. This cleavage results in separation of the 16S, the 23S and the 5S rRNA into precursor rRNA molecules (pre-rRNA molecules) which besides the rRNA species also contain leader and tail sequences. The primary RNase III cleavage is normally a rapid process, whereas the

subsequent maturation is substantially slower. Precursor rRNA is typically more abundant than even strongly expressed mRNA species. Thus, for certain applications, precursor rRNA may be an attractive diagnostic target. In order to specifically detect precursor rRNA, a target probe should be directed against sequences comprising at least part of the leader or tail sequences. A target probe may further be directed against sequences of which both part of the leader/tail and mature rRNA sequences are constituents.

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Usually, patients having contracted a mycobacterial infection are treated with medicaments until no mycobacteria can be found in the sputum. Except for culturing, the presently available methods do not allow for clear distinguishing between living and dead mycobacteria. This means that a patient may often be treated with medicaments for a longer period of time than actually necessary. A way of determining the progress of treatment would be a very valuable tool in the fight of tuberculosis and other mycobacterial diseases.

As transcription and maturation of rRNA is a measure of viability, detection of precursor rRNA is a suitable and direct measure of viability of the bacteria. Furthermore, precursor rRNA may be used for identification of antibiotic drugs which reduce or inhibit rRNA transcription. One such example is rifampicin. A transcriptional inhibitor will in susceptible bacteria eliminate new synthesis of rRNA and thus the pool of precursor rRNA will be depleted. However, in resistant cells, primary transcripts as well as precursor rRNAs will continue to be produced.

Although it is preferred to use peptide nucleic acid probes targeting specific sequences of rRNA, it will readily be understood that peptide nucleic acid probes complementary to rRNA targeting probes will be useful for the detection of the genes coding for said sequence specific rRNA (rDNA), and peptide nucleic acid probes for the detecting rDNA is hence contemplated by the present invention. Although it is preferred to choose the sequence of the probe so as to enable the probe to hybridise to its target sequence in antiparallel orientation, it is to be understood that probes capable of hybridising in parallel orientation can be constructed from the same information. The present invention is intended to cover both types of probes.

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In the broadest aspect, the present invention relates to peptide nucleic acid probes for detecting a target sequence of one or more mycobacteria optionally present in a test sample, said probe being capable of hybridising to a target sequence of mycobacterial rDNA, precursor rRNA or rRNA (claim 1).

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The probes of the invention may suitably be directed to rDNA, precursor rRNA, or to 23S, 16S or 5S rRNA.

In accordance with claim 3, the target sequences, to which the peptide nucleic acid probe(s) are capable of hybridising to, are obtainable by

- (a) comparing the nucleobase sequences of said mycobacterial rRNA or rDNA of one or more
 mycobacteria to be detected with the corresponding nucleobase sequence of organism(s), in particular other mycobacteria, from which said one or more mycobacteria are to be distinguished,
 - (b) selecting a target sequence of said rRNA or rDNA which includes at least one nucleobase differing from the corresponding nucleobase of the organism(s), in particular other mycobacteria, from which said one or more mycobacteria are to be distinguished, and
 (c) determining the capability of said probe to hybridise to the selected target sequence to form detectable hybrids.

Peptide nucleic acid probes are, in accordance with claim 4, obtainable by

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- (a) comparing the nucleobase sequences of said mycobacterial rRNA or rDNA of one or more mycobacteria to be detected with the corresponding nucleobase sequence of organism(s), in particular other mycobacteria, in particular other mycobacteria, from which said one or more mycobacteria are to be distinguished,
- (b) selecting a target sequence of said rRNA or rDNA which includes at least one nucleobase differing from the corresponding nucleobase of the organism(s), in particular other mycobacteria, from which said one or more mycobacteria are to be distinguished,
 (c) synthesising said probe, and
 - (4) determining the capability of said probe to hybridise to the selected target sequence to form detectable hybrids.

The probes are in particular suitable for detecting a target sequence of one or more mycobacteria of the Mycobacterium tuberculosis Complex (MTC) or for detecting a target sequence of one or more mycobacteria other than mycobacteria of the Mycobacterium tuberculosis Complex (MOTT) optionally present in a sample, which probe comprises from 6 to 30 polymerised peptide nucleic acid moieties, said probe being capable of hybridising to a target sequence of mycobacterial rDNA, precursor rRNA or 23S, 16S or 5S rRNA forming detectable hybrids (claim 5). In accordance with claim 6, such probes may comprise peptide nucleic acid moieties of formula (I)

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wherein each X and Y independently designate O or S, each Z independently designates O, S, NR^1 , or $C(R^1)_2$, wherein each R^1 independently designate H, C_{1-6} alkyl, C_{1-6} alkyl, C_{1-6} alkynyl,

each R², R³ and R⁴ designate independently H, the side chain of a naturally occurring amino acid, the side chain of a non-naturally occurring amino acid, C₁₋₄ alkyl, C₁₋₄ alkenyl or C₁₋₄ alkynyl, or a functional group, each Q independently designates a naturally occurring nucleobase, a non-naturally occurring nucleobase, an intercalator, a nucleobase-binding group, a label or H,

5 · and with the proviso indicated in claim 6.

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The probes may suitably be used for detecting a species specific mycobacterial target sequence, or target sequences of a group of mycobacteria like MTC, MOTT, MAC or MAIS. The probes may further be designed so as to be capable of hybridising to one or more drug resistant mycobacteria, or, alternatively, to the wild-type corresponding thereto. In the design of the probes, sequences between different mycobacteria (one or more) may be taken into account as may sequences from other related or non-related organisms (one or more).

As mentioned above, drug-resistance is an increasing threat to the fight of mycobacterial infection. Monotherapy with macrolides such as clarithromycin and azithromycin often leads to clinically significant drug-resistance. Clarithromycin and azithromycin are important drugs in the treatment of infections by especially M. avium. Comparison between 23S rRNA sequences from isolates of M. avium and M. intracellulare with acquired resistance to clarithromycin and azithromycin and 23S rRNA sequences from non-resistant strains has revealed that the majority of resistant strains have single-point mutations in the 23S rRNA in positions corresponding to 2058 and 2059 in E. coli 23S rRNA. In the M. avium 23S rRNA sequence (GenBank accession number X74494), the corresponding bases are in position 2568 and 2569, respectively, as shown in Figure 6. Most susceptible strains have an A residue in one of these positions whereas the resistant strains have a C, G or T in position 2058 (E. coli numbering corresponding to 2568 in M. avium with GenBank accession number X74494), or G or C in position 2059 (E. coli numbering corresponding to 2569 in M. avium with GenBank accession number X74494).

Single-point mutations in rRNA apparently also account to some degree for streptomycin resistance. Streptomycin, the first successful antibiotic drug against tuberculosis, is an aminocyclitol glycoside that perturbs protein synthesis at the ribosomal level. The genetic basis for streptomycin resistance has not yet been completely solved. However, some streptomycin resistant strains of M. tuberculosis have single-point mutations in 16S rRNA. These mutations are located in positions corresponding to bases 501, 522, 523, 526, 912 and 913 in E. coli 16S rRNA which correspond to bases with numbers 452, 473, 474, 477, 865 and 866, respectively, of M. tuberculosis 16S rRNA (GenBank accession number X52917) as shown in Figure 7. Most of these mutated nucleotides are involved in structural interactions within the 530 loop of 16S rRNA which is one of the most conserved regions in the entire 16S rRNA gene.

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Mutations in an 81 bp region of the gene (rpoB) encoding the beta subunit of RNA polymerase are the cause of 96% of the cases of rifampicin resistance in M. tuberculosis and M. leprae. rRNA precursor molecules have terminal domains (tails) which are removed during late steps in precursor rRNA processing to yield the mature rRNA molecules. Transcriptional inhibitors such as rifampicin can deplete precursor rRNA in sensitive cells by inhibiting de novo precursor rRNA synthesis while allowing maturation to proceed. Thus, precursor rRNA is depleted in susceptible mycobacterium cells while it remains produced in resistant mycobacterium cells when the cells are exposed to rifampicin during culturing.

Peptide nucleic acid probes for detecting a target sequence of one or more mycobacteria of the Mycobacterium tuberculosis Complex are defined in claims 7 to 10. Peptide nucleic acid probes for detecting a target sequence of one or more mycobacteria other than mycobacteria of the Mycobacterium tuberculosis Complex are defined in claims 11 to 13. Peptide nucleic acid probes for detecting a target sequence of one or more drug resistant mycobacteria of the Mycobacterium tuberculosis complex or of one or more drug resistant mycobacteria other than mycobacteria of the Mycobacterium tuberculosis Complex are defined in claim 14.

In the present context and the claims, the term "naturally occurring nucleobases" includes the four main DNA bases (i.e. thymine (T), cytosine (C), adenine (A) and guanine (G)) as well as other naturally occurring nucleobases (e.g. uracil (U) and hypoxanthine).

The term "non-naturally occurring nucleobases" comprises i.a. modified naturally occurring nucleobases. Such non-naturally occurring nucleobases may be modified by substitution by e.g. one or more C₁₋₈ alkyl, C₁₋₈ alkenyl or C₁₋₈ alkynyl groups or labels. Examples of non-naturally occurring nucleobases are purine, 2,6-diamino purine, 5-propynylcytosine (C propynyl), isocytosine (iso-C), 5-methyl-isocytosine (iso-C) (see e.g. Tetrahedron Letters Vol

36, No 12, 2033-2036 (1995) or Tetrahedron Letters Vol 36, No 21, 3601-3604 (1995)), 7-deazaadenine, 7-deazaguanine, N⁴-ethanocytosine, N⁶-ethano-2,6-diaminopurine, 5-(C₃₋₆)-alkenyluracil, 5-(C₃₋₆)-alkynylcytosine, 5-fluorouracil and pseudocytosine.

5 Examples of useful intercalators are e.g. acridin, antraquinone, psoralen and pyrene.

Examples of useful nucleobase-binding groups are e.g. groups containing cyclic or heterocyclic rings. Non-limiting examples are 3-nitro pyrrole and 5-nitro indole.

- It is to be understood that alkyl, alkenyl and alkynyl groups may be branched or non-branched, cyclic or non-cyclic, and may further be interrupted by one or more heteroatoms, or may be unsubtituted or substituted by or may contain one or more functional groups. Non-limiting examples of such functional groups are acetyl groups, acyl groups, amino groups, carbamido groups, carbamoyl groups, carbamyl groups, carbonyl groups, carboxy groups, cyano groups, dithio groups, formyl groups, guanidino groups, halogens, hydrazino groups, hydrazo groups, hydroxamino groups, hydroxy groups, keto groups, mercapto groups, nitro groups, phospho groups, phosphon ester groups, sulfo groups, thiocyanato groups, cyclic, aromatic and heterocyclic groups.
- C₁₋₄ groups contain from 1 to 4 carbon atoms, C₁₋₆ groups contain from 1 to 6 carbon atoms, and C₁₋₁₅ groups contain from 1 to 15 carbon atoms, not including optional substituents, heteroatoms and/or functional groups. Non-limiting examples of such groups are -CH₃, -CF₃, -CH₂-, -CH₂CH₃, -CH₂CH₂-, -CH(CH₃)₂, -OCH₃, -OCH₂-, -OCH₂CH₃, -OCH₂CH₂-, -OCH(CH₃)₂, -OC(O)CH₃, -C(O)CH₂-, -C(O)CH₃, -C(O)CH₃, -C(O)OH, -C(O)O-, -CH₂NH₂, -CH₂NH-, -CH₂OCH₃, -CH₂OCH₂-, -CH₂OC(O)OH, -CH₂OC(O)O-, -CH₂C(O)CH₂-, -C(O)NH₂, -CH=CH₂, -CH=CH-, -CH=CHCH₂C(O)OH, -CH=CHCH₂C(O)O-, -C=CH, -C=C-, -CH₂C=CH, -CH₂C=CCH₃, -OCH₂C=CCH, -OCH₂C=CCH₃, -NHCH₂C(O)-, -NHCH₂CH₂C(O)-, -NH(CH₂CH₂O)₂CH₂C(O)-, and HO(O)CCH₂C(O)(NH-(CH₂CH₂O)₂CH₂C(O))₂-, phenyl, benzyl, naphthyl, oxazolyl, pyridinyl, thiadiazolyl, triazolyl, and thienyl.

Within the present context, the expression "naturally occurring amino acid" is intended to comprise D- and L-forms of amino acids commonly found in nature, e.g. D- and L-forms of Ala (alanine), Arg (arginine), Asn (aspargine), Asp (aspartic acid), Cys (cysteine), Gln (glutamine), Glu (glutamic acid), His (histidine), Ile (isoleucine), Leu (leucine), Lys (lysine), Met (methionine), Phe (phenylalanine), Pro (proline), Ser (serine), Thr (threonine), Trp (tryptophan), Tyr (tyrosine) and Val (valine).

In the present context, the expression "non-naturally occurring amino acid" is intended to comprise D- and L-forms of amino acids other than those commonly found in nature as well as modified naturally occurring amino acids. Examples of useful non-naturally occurring amino acids are D- and L-forms of β -Ala (β -alanine) Cha (cyclohexylalanine), Cit (citrulline), Hci (homocitrulline), HomoCys (homocystein), Hse (homoserine), Nle (norleucine), Nva (norvaline), Orn (ornithine), Sar (sarcosine) and Thi (thienylalanine).

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In the present context, the term "sample" is intended to cover all types of samples suitable for the purpose of the invention. Examples of such samples are sputum, laryngeal swabs, gastric lavage, bronchial washings, biopsies, aspirates, expectorates, body fluids (spinal, pleural, pericardial, synovial, blood, pus, bone marrow), urine, tissue sections as well as food samples, soil, air and water samples. Analysis of samples originating from the before-mentioned samples (e.g. cultures and treated samples) are also within the scope of the invention.

In the present context, the term "hybrids" is intended to include complexes between a probe and a nucleic acid to be determined. Such hybrids may be made up of two or more strands.

The strength of the binding between the probe and the target nucleic acid sequence may be influenced by the ligand Q. When Q designates a nucleobase, Hoogsteen and/or Watson-Crick base pairing assist(s) in the formation of hybrids between a nucleic acid sequence to be detected and a probe. It is contemplated that one or more of the ligands may be a group which contribute little or none to the binding of the nucleic acid such as hydrogen. It is contemplated that suitable probes to be used comprise less than 25% by weight of peptide nucleic acid moieties, wherein Q designates such groups. One or more of the ligands Q may be groups that stabilise nucleobase stacking such as intercalators or nucleobase-binding groups.

In the above-indicated probes, one or more of the Q-groups may designate a label. Examples of suitable labels are given below. Moieties wherein Q denotes a label may preferably be located in one or both of the terminating moieties of the probe. Moieties wherein Q denotes a label may, however, also be located internally.

The peptide nucleic acid probes may comprise moieties, wherein all X groups are O (polyamides) or wherein all X groups are S (polythioamides). It is to be understood that the probes may also comprise mixed moieties (comprising both amide and thioamide moieties).

In another aspect, the present invention relates to peptide nucleic acid probes of formula (II), (III) and (IV) as well as mixtures of such probes defined in claim 15.

In a preferred embodiment, the peptide nucleic acid probes or mixtures thereof according to the invention are of formulas (I)-(IV) as defined in claim 16 with Z being NH, NCH₃ or O, each R^2 , R^3 and R^4 independently being H or the side chain of a naturally occurring amino acid, the side chain of a non-naturally occurring amino acid, or C_{1-4} alkyl, and each Q being a naturally occurring nucleobase or a non-naturally occurring nucleobase with the provisos defined in claims 6 to 14.

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Peptide nucleic acid probes or mixtures of such probes according to the invention are preferably those of formula (I)-(IV) as defined in claim 17 with Z being NH or O, and R² being H or the side chain of Ala, Asp, Cys, Glu, His, HomoCys, Lys, Orn, Ser or Thr, and Q being a nucleobase selected from thymine, adenine, cytosine, guanine, uracil, iso-C, and 2,6-diaminopurine with the provisos defined in claims 6 to 14.

Peptide nucleic acid probes or mixtures thereof, which are of major interest for detecting mycobacteria of the MTC group or one or more mycobacteria other than mycobacteria of the MTC group, are probes of formula (V) according to claim 18, wherein R⁴ is H or the side chain of Ala, Asp, Cys, Glu, His, HomoCys, Lys, Orn, Ser or Thr, Q is as defined in claim 17 and with the provisos indicated in claims 6 to 14.

The peptide nucleic acid probe comprises polymerised moieties as defined above and in the claims. From the formula, it is to be understood that the probe may comprise polymerised moieties which structure may be mutually different or identical. In some cases, it may be advantageous that at least one moiety of the probe, preferably one (or both) of the moieties terminating the probe, are of a different structure. Such terminating moieties may suitably be a moiety of formula (VI)

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where Q is as defined above. Such moiety may suitably be connected to a peptide nucleic acid moiety through an amide bond.

The peptide nucleic acid probe according to the invention comprises from 6 to 30 polymerised moleties of formulas (I) to (V), and, in addition, optionally one or two terminating moleties of formula (VI) as defined above. The preferred length of the probe will depend on the sample material and whether labelled probes are used. It is contemplated that especially interesting probes comprise from 10 to 30 polymerised moleties of formulas (I) to (V), and, in addition,

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optionally one or two terminating moieties of formula (VI) as defined above. Probes of the invention may suitably comprise from 12 to 25 polymerised moieties of formulas (I) to (V), more suitably from 14 to 22 polymerised moieties of formulas (I) to (V), most suitably from 15 to 20 polymerised moieties of formulas (I) to (V), and, in addition, optionally one or two terminating moieties of formula (VI).

As mentioned above, the polymerised moieties of the probes may be mutually different or identical. In some embodiments, the polymerised moieties of formulas (V) constitute at least 75% by weight (calculated by excluding labels and linkers), preferably at least 80% by weight and most preferably at least 90% by weight of the probe.

The ends on the moieties terminating the probe may be substituted by suitable substituents which in the following will be named "linkers". A terminating end may suitably be substituted by from 1 to 5 linkers, more suitably from 1 to 3 linkers. Such linkers may suitably be selected among C₁₋₁₅ alkyl, C₁₋₁₅ alkenyl and C₁₋₁₅ alkynyl groups as defined above. The linkers may be substituted or unsubstituted, branched or non-branched, or be interrupted by heteroatoms, or be substituted or contain functional groups as described above. This may depend on the chemical nature of the terminating moiety (i.e. whether the moiety is terminated by a carbon, oxygen or nitrogen atom). A terminating end or a linker on a terminating end may further be substituted by one or more labels, which labels may be incorporated end to end, i.e. so as to form a non-branched labelled end, or may be incorporated so as to form a branched labelled end ("zipper"). The linkers may be attached directly to a terminating end, may be attached to a label or between labels on a terminating end, or be attached to a terminating end before a label is attached to a terminating end. It should be understood that two terminating ends may carry different or identical substituents, linkers and/or labels. It should further be understood that the term "a label" is intended to comprise one or more labels as the term "linkers" is to comprise one or more linkers. For certain applications, it may be advantageous that one or more linkers are incorporated between the peptide nucleic acid moieties. Such applications may in particular be those based on triplex formation.

Examples of suitable linkers are -NH(CH₂CH₂O)_nCH₂C(O)-, -NH(CHOH)_nC(O)-, -(O)C(CH₂OCH₂)_nC(O)- and -NH(CH₂)_nC(O)-, NH₂(CH₂CH₂O)_nCH₂C(O)-, NH₂(CHOH)_nC(O)-, HO(O)C(CH₂OCH₂)_nC(O)-, NH₂(CH₂)_nC(O)-, -NH(CH₂CH₂O)_nCH₂C(O)OH, -NH(CHOH)_nC(O)OH, -(O)C(CH₂OCH₂)_nC(O)OH and -NH(CH₂)_nC(O)OH, wherein n is 0 or an integer from 1 to 8, preferably from 1 to 3. Examples of very interesting linkers are

-NHCH₂C(O)-, -NHCH₂CH₂C(O)-, -NH(CH₂CH₂O)₂CH₂C(O)-, and HO(O)CCH₂CH₂C(O)(NH(CH₂CH₂O)₂CH₂C(O))₂-.

In the present context, the term "label" refers to a substituent which is useful for detection or visualisation. Suitable labels comprise fluorophores, biotin, dinitro benzoic acid, digoxigenin, radioisotope labels, peptide or enzyme labels, chemiluminiscence labels, fluorescent particles, hapten, antigen or antibody labels.

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The expression "peptide label" is intended to mean a label comprising from 1 to 20 naturally occurring or non-naturally occurring amino acids, preferably from 1 to 10 naturally occurring or non-naturally occurring amino acids, more preferably from 1 to 8 naturally occurring or non-naturally occurring amino acids, most preferably from 1 to 4 naturally occurring or non-naturally occurring amino acids, linked together end to end in a non-branched or branched ("zipper") fashion. Such peptide label may be composed of amino acids which are mutually identical or different. In a preferred embodiment, such a non-branched or branched end comprises one or more, preferably from 1 to 8 labels, more preferably from 1 to 4, most preferably 1 or 2, further labels other than a peptide label. Such further labels may suitably terminate a non-branched end or a branched end. One or more linkers may suitably be attached to the terminating end before a peptide label and/or a further label. Furthermore, such peptide labels may be incorporated between the peptide nucleic acid moieties.

The probe as such may also comprise one or more labels such as from 1 to 8, preferably from 1 to 4, most preferably 1 or 2, labels and/or one or more linker units, which may be attached internally, i.e. to the backbone of the probe. The linker units and labels may mutually be attached as described above.

Examples of particular interesting labels are biotin, fluorescein labels, e.g. 5-(and 6)-carboxy-fluorescein, 5- or 6-carboxyfluorescein, 6-(fluorescein)-5-(and 6)-carboxamido hexanoic acid and fluorescein isothiocyanate, peptide labels consisting of from 1 to 20 naturally occurring amino acids or non-naturally occurring amino acids, enzyme labels such as peroxidases like horse radish peroxidase (HRP), alkaline phosphatase, and soya bean peroxidase, dinitro benzoic acid, rhodamine, tetramethylrhodamine, cyanine dyes such as Cy2, Cy3 and Cy5, coumarin, R-phycoerythrin (RPE), allophycoerythrin, Texas Red, Princeton Red, and Oregon Green as well as conjugates of R-phycoerythrin and, e.g. Cy5 or Texas Red.

Examples of preferred labels are biotin, fluorescent labels, peptide labels, enzyme labels and dinitro benzoic acid. Peptide labels may preferably be composed of from 1 to 10, more preferably of from 1 to 8, most preferably of from 1 to 4, naturally occurring or non-naturally occurring amino acids. It may be particularly advantageous to incorporate one or more other labels as well as a peptide label such as from 1 to 8 or from 1 to 4 other labels, preferably 1 or

2 other labels.

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Suitable peptide labels may preferably be composed of cysteine, glycine, lysine or ornithine.

- In a further embodiment, the Q substituent as defined above may be labelled. Suitable labels are as defined above. Between Q and such a label, a linker as defined above may be incorporated. It is preferred that such labelled ligands Q are selected from thymine and uracil labelled in the 5-position and 7-deazaguanine and 7-deazaguanine labelled in the 7-position.
- A mixture of peptide nucleic acid probes is also part of the present invention. Such mixture may comprise more than one probe capable of hybridising to 23S rRNA, and/or more than one probe capable of hybridising to 16S rRNA, and/or or more than one probe capable of hybridising to 5S rRNA. A mixture of probes may further comprise probe(s) directed to precursor rRNA and/or rDNA. The mixture may also comprise peptide nucleic acids for detecting more than one mycobacteria in the same assay.
 - In a preferred embodiment, the nucleobase sequence of the peptide nucleic acid probe is selected so as to be substantially complementary to the nucleobase sequence of the target sequence in question. In an especially preferred embodiment, the nucleobase sequence of the peptide nucleic acid probe is selected so as to be complementary to the nucleobase sequence of the target sequence in question. By "complementary" is meant that the nucleobases are selected so as to enable perfect match between the nucleobases of the probe and the nucleobases of the target, i.e. A to T or G to C. By substantially complementary is meant that the peptide nucleic acid probe is capable of hybridising to the target sequence, however, the probe does not necessarily have to be perfectly complementary to the target. For example, probes comprising one or more bases not complementary to the target sequence and nontarget sequences, especially base(s) located at the end of the probe, where the effect on the stability of probe-target nucleic acid hybrids is low. Another example is probes comprising other naturally occurring bases. Thus provided that the probe is capable of hybridising to the target sequence, the nucleobase difference(s) between target sequences and non-target sequences ensures that the stability of probe-non-target nucleic acid hybrids are lower than the stability of probe-target nucleic acid hybrids and therefore make such substantially complementary probes applicable for detection of mycobacteria.
- The probes may be synthesised according to the procedures described in "PNA Information Package" obtained from Millipore Corporation (Bedford, MA, USA), or may be synthesised on an Expedite Nucleic Acid Synthesis System (PerSeptive BioSystems, USA).

If using the Fmoc strategy for elongation of the probe with linkers or amino acids, it is possible to retain side chain amino groups protected with acid sensitive protection groups such as the Boc or Mtt group. This method allows introduction of a linker containing several Boc protected amino groups which can all be cleaved and labelled in the same synthesis cycle.

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One way of labelling a probe is to use a fluorescent label, such as 5-(and 6)-carboxyfluorescein, 5- or 6-carboxyfluorescein, or 6-(fluorescein)-5-(and 6)-carboxamido hexanolc acid. The acid group is activated with HATU (O-(7-azabenzotriazol-1-yl)-1,1,3,3-tetramethyluronium hexafluorophosphate) or HBTU (2-(1H-benzotriazol-1-yl)-1,1,3,3-tetramethyluronium hexafluorophosphate) and reacted with the N-terminal amino group of the peptide nucleic acid. The same technique can be applied to other labelling groups containing an acid function. Alternatively, the succinimidyl ester of the above-mentioned labels may suitably be used or fluorescein isothiocyanate may be used directly.

After synthesis, probes can be cleaved from the resin using standard procedures as described by Millipore Corporation or PerSeptive BioSystems. The probes are subsequently purified and analysed using reversed-phase HPLC techniques at 50°C and were characterised by matrix-assisted laser desorption/ionisation time of flight mass spectrometry (MALDI-TOFMS), plasma desorption mass spectrometry (PDMS), electron spray mass spectrometry (ESMS), or fast atom bombardment (FAB-MS).

Generally, probes such as probes comprising polymerised moieties of formula (IV) and (V) may also be prepared as described in, e.g., Angewandte Chemie, International Edition in English 35, 1939-1942 (1996) and Bioorganic & Medical Chemistry Letters, Vol 4, No 8, 1077-1080 (1994). Chemical properties of some probes are described in, e.g., Nature, 365, 566-568 (1993).

The method as claimed can be used for the detection of a target sequence of one or more mycobacteria optionally present in a sample. The method and the probes provide a valuable tool for analysing samples for the presence of such target sequences, hence providing information for establishing a diagnosis.

In the assay method according to the invention, the sample to be analysed for the presence of mycobacteria is brought into contact with one or more probes or a mixture of such probes according to the invention under conditions by which hybridisation between the probe(s) and any sample rRNA or rDNA originating from mycobacteria can occur, and the formed hybrids, if any, are observed or measured, and the observation or measurement is related to the presence of a target sequence of one or more mycobacteria. The observation or

Prior to contact with probe(s) according to the invention, the samples may undergo various

measurement may be accomplished visually or by means of instrumentation.

types of sample processing which include purification, decontamination and/or concentration.

The sample may suitably be decontaminated by treatment with sodium hypochlorite and subsequently centrifuged for concentration of the mycobacteria. Samples e.g. sputum samples may be treated with a mucolytic agent such as N-Acetyl-L-cystein which reduces the viscosity of the sample as well as be treated with sodium hydroxide which decontaminates the sample, and subsequently centrifuged. Other well-known decontamination and concentration procedures include the Zephiran-trisodium phosphate method, Petroff's sodium hydroxide method, the oxalic acid method as well as the cetylpyridinium chloride-sodium chloride method. Samples may also be purified and concentrated by applying sample preparation methods such as filtration, immunocapture, two-phase separation either alone or in

combination. The sample preparation methods may also be used together with the

centrifugation and decontamination methods mentioned above.

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Samples may, either directly or after having undergone one or more processing steps, be analysed in primarily two major types of assays, in situ-based assays and in vitro-based assays. In this context, in situ-based assays are to be understood as assays, in which the target nucleic acids remain within the bacterial cell during the hybridisation process. Examples are in situ hybridisation (ISH) assays on smears and biopsies as well as hybridisation to whole cells which may be in suspension and which subsequently may be analysed by e.g. flow cytometry optionally after capture of the bacteria onto particles (with same or different type and size), or by image analysis after spreading of the bacteria onto a solid medium, filter membrane or another substantially planar surface.

In vitro-based assays are to be understood as assays, in which the target nucleic acids are released from the bacterial cell before hybridisation. Examples of such assays are microtiter plate-based assays. Many other assay types, in which the released target nucleic acids by some means are captured onto a solid phase and subsequently analysed via a detector probe, are feasible and within the scope of the present invention. Even further, in vitro-based assays include assays, in which the target nucleic acids are not captured onto a solid phase, but in which the hybridisation and signal generation take place entirely in solution.

Samples for in situ-based assays may suitably be applied and optionally be immobilised to a support. Techniques for applying of a sample onto a solid support depend on the type of sample in question and include smearing and cytocentrifugation for liquid or liquified samples and sectioning of tissues for biopsy materials. The solid support may take a wide variety of

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forms well-known in the art, such as a microscope slide, a filter membrane, a polymer membrane or a plate of various materials.

In the case of in vitro-based assays, the target nucleic acid may be released from the mycobacterial cells in various ways. Most methods for releasing the nucleic acids cause bursting of the cell wall (lysis) followed by extraction of the nucleic acids into a buffered solution. As mycobacteria have complex cell walls containing covalently associated peptidoglycans, arabinogalactans and in particular mycolic acids, they cannot easily be disrupted by standard methods used for the rapid lysis of other bacteria. Possible methods which are known to give successful lysis of the mycobacterial cell wall include methods which involve treatment with organic solvents, treatment with strong chaotropic reagents such as high concentrations of guanidine thiocyanate, enzyme treatment, bead beating, heat treatment, sonication and/or application of a French press.

Samples to be analysed by in situ assays may be fixed prior to hybridisation. The person skilled in the art will readily recognise that the appropriate procedure will depend on the type of sample to be examined. Fixation and/or immobilisation should preferably preserve the morphological integrity of the cellular matrix and of the nucleic acids. Examples of methods for fixation are flame fixation, heat fixation, chemical fixation and freezing. Flame fixation may be accomplished by passing the slide through the blue cone of a Bunsen burner 3 or 4 times; heat fixation may be accomplished by heating the sample to 80°C for 2 hours; chemical fixation may be accomplished by immersion of the sample in a fixative (e.g. formamide, methanol or ethanol). Freezing is particularly relevant for biopsies and tissue sections and is usually carried out in liquid nitrogen.

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In one in situ hybridisation assay embodiment, the sample to be analysed is smeared onto a substantially planar solid support which may be a microscope slide, a filter membrane, a polymer membrane or another type of solid support with a planar surface. The preferred solid support is a microscope slide. After the smear has been prepared, it may optionally undergo further pre-treatment like treatment with bactericidal agents or additional fixation by immersion in e.g. ethanol. The sample may also be pre-treated with enzyme(s) which as primary function permeabilise the cells and/or reduce the viscosity of the sample. It may further be advantageous to perform a pre-hybridisation step in order to block sites which might otherwise give raise to non-specific binding. For this purpose, blocking agents like skim milk, and non-target probes may suitably be used. The components of the pre-hybridisation mixture should be selected so as to obtain an effective saturation of sites in the sample that might otherwise bind the probe non-specifically. The pre-hybridisation buffer may suitably comprise an appropriate buffer, blocking agent(s), and detergents.

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During the in situ hybridisation, one or more probes according to the present invention are brought into contact with any target rRNA or rDNA inside the cells in a hybridisation solution under suitable stringency conditions. The concentration of the applied probe may vary depending on the chemical nature of the probe and the conditions under which hybridisation is carried out. Typically, a probe concentration between 1 nM and 1 µM is suitable. The hybridisation solution may comprise a denaturing agent which allows hybridisation to take place at a lower temperature than would be the case without the agent. The denaturing agent should be present in an amount effective to increase the ratio between specific binding and non-specific binding. The effective amount of denaturing agent depends on the type used and on the probe or combination of probes. Examples of denaturing agents are formamide, ethylene glycol and glycerol, and these may preferably be used in a concentration above 10% and less than 70%. The preferred denaturing agent is formamide which is used more preferably in concentrations from 20% to 60%, most preferably from 30% to 50%. It should be noted that in several instances it may not be necessary or advantageous to include a denaturing agent.

Prior to hybridisation or during hybridisation, a mixture of random probes (probes with random, non-selected sequences of optionally different length) may be added in excess to reduce non-specific binding. Also, one or more non-sense probes (probes with a defined nucleobase sequence and length differing from the nucleobase sequence of the target sequence) may be added in excess in order to reduce non-specific binding. Also, non-specific binding of detectable probes to one or more non-target nucleic acid sequences can be suppressed by addition of one or more unlabelled or independently detectable probes, which probes have a sequence that is complementary to the non-target sequence(s). It is particularly advantageous to add such blocking probes when the non-target sequence differs from the target sequence by only one mismatch.

It may be advantageous to include inert polymers which are believed to increase the effective concentration of the probe(s) in the hybridisation solution. One such macromolecule is dextran sulphate which may be used in concentrations of from 2.5% to 15%. The preferred concentration range is from 8% to 12% in the case of dextran sulphate. Other useful macromolecules are polyvinylpyrrolidone and ficoll, which typically are used at lower concentrations, e.g. 0.2%. It may further be advantageous to add one or more detergents which may reduce the degree of non-specific binding of the peptide nucleic acid probes. Examples of useful detergents are sodium dodecyl sulphate, Tween 20® or Triton X-100®. Detergents are usually used in concentrations between 0.05% and 1.0%, preferably between 0.05% and 0.25%. The hybridisation solution may furthermore contain salt. Although it is not

necessary to include salt in order to obtain proper hybridisation, it may be advantageous to include salt in concentrations from 2 to 500 mM, or suitably from 5 to 100 mM.

During hybridisation, other important parameters are hybridisation temperature, concentration of the probe and hybridisation time. The person skilled in the art will readily recognise that optimal conditions must be determined for each of the above-mentioned parameters according to the specific situation, e.g. choice of probe(s) and type and concentration of the components of the hybridisation buffer, in particular the concentration of denaturing agent. Presence of volume excluders may also have an effect.

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Following hybridisation, the sample is washed to remove any unbound and any non-specifically bound probe, and consequently, appropriate stringency conditions should be used. By stringency is meant the degree to which the reaction conditions favour the dissociation of the formed hybrids. The stringency may be increased typically by increasing the washing temperature and/or washing time. Typically, washing times from 5 to 40 minutes may be sufficient. Two or more washing periods of shorter time may also give good results. A range of buffers may be used, including biological buffers, phosphate buffers and standard citrate buffers. The demand for low salt concentration in the buffers is not as pertinent as for DNA probe assays due to the difference response to salt concentration. In some cases, it is advantageous to increase the pH of the washing buffer as it may give an increased signal-to noise ratio (see WO 97/18325). This is conceivably due to a significant reduction of the non-specific binding. Thus, it may be advantageous to use a washing solution with a pH value form 8 to 10.5, preferably from 9 to 10.

Visualisation of bound probe(s) must be carried out with due regard to the type of label chosen. There are a wide range of useful probe labels, in particular various fluorescent labels such as fluorescein, rhodamine and derivatives thereof. Furthermore, labels like enzymes (e.g. peroxidases and phosphatases) and haptens (e.g. biotin, digoxigenin, dinitro benzoic acid) may suitably be applied. In the case of fluorescent labels, the hybrids formed may be visualised using a microscope with a magnification of at least × 250, preferably × 1000. The visualisation may further be carried out using a CCD (charge coupled device) camera optionally controlled by a computer. When haptens are used as labels, the hybrids may be detected using an antibody conjugated with an enzyme. In these cases, the detection step may be based on colorimetry, fluorescence or luminescence.

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The probes may alternatively be labelled with fluorescent particles having the fluorescent label embedded in the particles (e.g. Estapor K coloured microspheres), located on the surface of the particles and/or coupled to the surfaces of the particles. As the particles have to come into

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necessitate pretreatment of the bacteria. Relatively small particles e.g. about 20 nm may suitable be used.

In another in situ hybridisation embodiment, frozen tissue or biopsy samples are cut into thin sections and transferred to a substantially planar surface, preferably microscope slides. Prior to hybridisation, the tissue or biopsy may be treated with a fixative, preferably a precipitating fixative such as acetone, or the sample is incubated in a solution of buffered formaldehyde. Alternatively, the biopsy or tissue section can be transferred to a fixative such as buffered formaldehyde for 12 to 24 hours and following fixation, the tissue may be embedded in paraffin forming a block from which thin sections can be cut. Prior to hybridisation, the tissue section is dewaxed and rehydrated using standard procedures. Permeabilisation (e.g. treatment with proteases, diluted acids, detergents, alcohol and/or heat) may in some cases be advantageous. The selected method for permeabilisation depends on several factors, for instance on the fixative used, the extent of fixation, the type and size of sample, and on the applied probe. For these types of samples, sample processing, prehybridisation, hybridisation, washing and visualisation may be carried out using same or adjusted conditions as described above.

In a further embodiment of the in situ assays, the bacterial cells are kept in suspension during fixation, prehybridisation, hybridisation and washing are carried out under the same or similar conditions as described above. The preferred type of label for this embodiment is fluorescent labels. This allows detection of hybridised cells by flow cytometry, recording the intensity of fluorescence per cell. Bacterial cells in suspension may further be coupled to particles, preferably with a size of from 20 nm to 10 µm. The particles may be made of materials wellknown in the art like latex, dextran, cellulose and/or agarose, and may optionally be paramagnetic or contain a fluorescent label. Normally, bacterial cells are coupled to particles using antibodies against the target bacteria, but other means like molecular imprinting may also be used. Coupling of the bacterial cells to particles may be advantageous in sample handling and/or during detection.

In the embodiments of in situ hybridisation described above, the probes according to the invention are used for detecting a target sequence of one or more mycobacteria. In a preferred embodiment, the probes are suitable for detecting a target sequence of mycobacteria of the Mycobacterium tuberculosis Complex (MTC), mycobacteria other than the Mycobacterium tuberculosis Complex (MOTT), or mycobacteria of the Mycobacterium avium Complex (MAC). The probes are further suitable for detecting simultaneously different target sequences originating from the same mycobacteria.

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Samples to be analysed using in vitro-based assays need to undergo a treatment by which the nucleic acids are released from the bacterial cells. Nucleic acids may be released using organic solvents, strong chaotropic reagents such as high concentrations of guanidine thiocyanate, enzymes, bead beating, heating, sonication and/or application of a French press. The obtained nucleic acids may undergo additional purification prior to hybridisation.

In one in vitro hybridisation embodiment, the sample comprising the target nucleic acid is added to a container comprising immobilised capture probe(s) and one or more probe(s) labelled to function as detector probe(s). The hybridisation should be performed under suitable stringency conditions. The hybridisation solution may further comprise a denaturing agent, blocking probes, inert polymers, detergents and salt as described for the in situ-type assays. Likewise, the hybridisation temperature, probe concentration and hybridisation time are important parameters that need to be controlled according to the specific conditions of the assay, e.g. choice of peptide nucleic acid probe(s) and concentration of some of the ingredients of the hybridisation buffer. If hybridisation of the target nucleic acid to the capture probe(s) and detector probe(s), respectively, is performed in two separate steps, different parameters, in particular different stringency conditions, may be used in these steps. The concentration of the capture probe may be higher for in situ assays as hybridisation may be controlled better and washing can be performed more efficiently.

The capture probes may be immobilised onto a solid support by any means, e.g. by a coupling reaction between a carboxylic acid on a linker and an amino derivatised support. The capture probe may further be coupled onto the solid support by photochemical activation of photoreactive groups which have been attached absorptively to the solid support prior to photochemical activation. Such photoreactive groups are described in the US 5 316 784 A. The capture probes may further be coupled to a hapten which allows an affinity based immobilisation to the solid support. One such example is coupling of a biotin to the probe(s) and immobilisation via binding to a steptavidin-coated surface.

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The solid support may take a wide variety of forms well-known in the art, such as a microtiter plate having one or more wells, a filter membrane, a polymer membrane, a tube, a dip stick, a strip and particles. Filter membranes may be made of cellulose, celluloseacetate, polyvinylidene fluoride or any other materials well-known in the art. The polymer membranes may be of polystyrene, nylon, polypropylene or any other materials well known in the art. Particles may be paramagnetic beads, beads made of polystyrene, polypropylene, polyethylene, dextran, nylon, amyloses, celluloses, polyacrylamides and agarose. When the solid support has the form of a filter, a membrane, a strip or beads, it (they) may be

incorporated into a single-use device.

The selection of the label of the detector probe(s) depend on the specific assay format and possible instrumentation. When biotin labelled probes are used, the hybrids may be detected using streptavidin or an antibody against the biotin label which antibody or streptavidin may be conjugated with an enzyme and the actual detection depend on the choice of the specific enzyme, preferably a phosphatase or a peroxidase, and the substrate for the selected enzyme. The signal may in some cases be enhanced using commercially available amplification systems such as the catalysed signal amplification system for biotinylates probes (CSA by DAKO). Various polymer-based enhancement systems may also be used. An example is a dextran polymer to which both a hapten specific antibody and an enzyme is coupled. The detector probe(s) may further be labelled with other haptens, e.g. digoxigenin. dinitro benzoic acid and fluorescein, in which case the hybrids may be detected using an antibody against the hapten which antibody may be conjugated with an enzyme. It is even possible to apply detector probe(s) which have enzymes coupled directly onto the probes. There are a wide range of possibilities for selection of enzyme substrates allowing for colourimetric (substrates e.g. p-nitro-phenyl phosphate or tetra-methyl-benzidine), fluorogenic (substrates e.g. 4-methylumbilliferylphosphate) or chemiluminescent (substrates e.g. 1,2dioxetanes) detection.

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The detector probes may further be labelled with various fluorescent labels, preferably fluorescein or modamine, in which case the hybrids may be detected by measuring the fluorescence.

The detector probe(s) will typically be different from the capture probe(s), thus ensuring dual species specificity. The dual specificity will most often allow at least one of the probes to be shorter, e.g. a 10 mer probe.

Furthermore, the capture of purine rich sequences may be improved by utilising bis-peptide nucleic acids as capture probes. Such bis-peptide nucleic acids are described in WO 96/02558. The bis-peptide nucleic acids comprise a first peptide nucleic acid strand capable of hybridising in parallel fashion to the target nucleic acid, and a second peptide nucleic acid strand capable of hybridising in antiparallel fashion to the purine rich sequence of the nucleic acid to be captured. The two peptide nucleic acid strands are connected by a linker and are in this way capable of forming a triplex structure with said purine rich sequence nucleic acid. The number of polymerised moieties of each linker-separated peptide nucleic acid may be as previously defined for non-bis-peptide nucleic acids. However, due to the high stability of the triplexes formed, bis-peptide nucleic acids with short first and second strands can be used

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making the design of a pyrimidine rich probe easier.

Instead of using a detector probe, capture probe: nucleic acid complexes may be detected using a detection system based on an antibody reacting specifically with complexes formed between peptide nucleic acids and nucleic acids (such as described in WO 95/17430), in which detection system the primary antibody may comprise a label, or which detection system comprises a labelled secondary antibody, which specifically binds to the primary antibody. The specific detection again depends on the selected substrate which may be of any type of those mentioned above.

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Depending on the type of specific assay format, label and detection principle various types of instrumentation may be used including conventional microplate readers, luminometers and flow cytometers. Adaptation of adequate instrumentation may allow for automatisation of the assay.

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In an example of this embodiment, a capture probe of the present invention is coupled to a microtiter plate by a photochemical reaction between antraquinon-labelled capture probe and polystyrene of the microwell. Target rRNA is added to the microwells and incubated under stringent conditions. Unbound rRNA is removed by washing and the microwell are incubated with a hapten-labelled detector probe under stringent conditions. The visualisation is carried out using an enzyme-labelled antibody against the hapten, which after removal of unbound antibody is detected using a chemiluminescence substrate.

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In another example of this embodiment capture probes are coupled to latex particles, and hybridisation is carried out under suitable conditions in the presence of e.g. fluorescein labelled detector probe(s). After hybridisation and optionally washing, the hybrids are detected by flow cytometry. A range of different beads (e.g. by size or colours) may carry different capture probes for different targets, thus allowing a multiple detection system.

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In a further embodiment of the in vitro assays format, the capture probe, the target nucleic acid and the detector probe may hybridise in solution, and subsequently the capture probe is attached to a solid phase. The solid phase, the hybridisation conditions and means of detection may be selected according to the specific method as described above.

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In a further embodiment of in vitro assays, the target nucleic acid may be immobilised onto filter or polymer membranes or other types of solid phases well-known in the art. The hybridisation conditions and means of detection may be selected according to the specific setup as described above.

In a further embodiment of the in vitro assay, an array of up to 100 or even more different probes directed against different target sequences may be immobilised onto a solid surface and hybridisation of the target sequences to all the probes is carried out simultaneously. The solid phase, the hybridisation conditions and means of detection may be as described above. This allow for simultaneous detection or identification of a range of parameters, i.e. species identification and resistance patterns.

The present probes further provide a method of diagnosing infection by mycobacteria and a method for determining the stage of the infection and the appropriate treatment by which methods one or more optionally labelled probes according to the invention are brought into contact with a patient sample and the type of treatment and/or the effect of a treatment is (are) evaluated.

Kits comprising at least one peptide nucleic acid probe as defined herein are also part of the present invention. Such kit may further comprise a detection system with at least one detecting reagent and/or a solid phase capture system.

DESCRIPTION OF SPECIFIC EMBODIMENTS

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Examples of suitable Qs of adjacent moieties are given below. Peptide nucleic acid probes comprising such Qs will be suitable for detecting mycobacteria, in particular mycobacteria of the MTC group or mycobacteria other than mycobacteria of the MTC group. The probes are written from left to right corresponding to from the N-terminal end towards the C-terminal end. Suitable Q subsequences for detecting 23S and 16S rRNA as well as 5S rRNA of the MTC group are given below. Suitable Q subsequences for detecting 23S and 16S rRNA of mycobacteria other than mycobacteria of the MTC group are further given below. The Q subsequences include at least one nucleobase complementary to a nucleobase selected from the positions given in parenthesis. The Q subsequences are given as non-limiting examples of construction of suitable probe nucleobase sequences. It is to be understood that the probes may comprise fewer or more peptide nucleic acid moieties than indicated.

MTC group (23S)

	AGA TGC GGG TAG CAC (selected from positions 149-158 in Figure 1A),	(Seq ID no 1)
35	TGT TTT CTC CTC CTA (selected from positions 220-221 in Figure 1A),	(Seq ID no 2)
	ACT GCC TCT CAG CCG (selected from positions 328-361 in	
	Figure 1A and Figure 1B),	(Seq ID no 3)
	TGA TAC TAG GCA GGT (selected from positions 453-455 in Figure 1B),	(Seq ID no 4)
	CGG ATT CAC AGC GGA (selected from positions 490-501 in Figure 1B),	(Seq ID no 5)

	TCA CCA CCC TCC TCC (calcated from next)	
	TCA CCA CCC TCC TCC (selected from positions 637-660 in Figure 1C),	(Seq ID no 6)
	CCA CCC TCC TCC (selected from positions 637-660 in Figure 1C)	(modified Seq ID no 6)
	TTA ACC TTG CGA CAT (selected from positions 706-712 in Figure 1D),	(Seq ID no 7)
_	ACT ATT CAC ACG CGC (selected from positions 762-789 in Figure 1D),	(Seq ID no 8)
5	CTC CGC GGT GAA CCA (selected from position 989 In Figure 1D),	(Seq ID no 9)
	GCT TTA CAC CAC GGC (selected from positions 1068-1072 in Figure 1E),	(Seq ID no 10)
	ACG CTT GGG GGC CTT (selected from position 1148 in Figure 1E),	(Seq ID no 11)
	CCA CAC CCA CAA (selected from positions 1311-1329 in Figure 1E),	(Seq ID no 12)
	CCG GTG GCT TCG CTG (selected from positions 1361-1364 in Figure 1F),	(Seq ID no 13)
10	ACT TGC CTT GTC GCT (selected from position 1418 in Figure 1F),	(Seq ID no 14)
	GAT TCG TCA CGG GCG (selected from positions 1563-1570 in Figure 1F),	(Seq ID no 15)
	AAC TCC ACA CCC CCG (selected from positions 1627-1638 in Figure 1G),	(Seq ID no 16)
	ACT CCA CAC CCC CGA (selected from positions 1627-1638 in Figure 1G),	(Seq ID no 17)
	ACC CCT TCG CTT GAC (selected from positions 1675-1677 in Figure 1G),	(Seq ID no 18)
15	CTT GCC CCA GTG TTA (selected from position 1718 in Figure 1G),	(Seq ID no 19)
	CTC TCC CTA CCG GCT (selected from positions 1734-1740 in Figure 1H),	(Seq ID no 20)
	GAT ATT CCG GTC CCC (selected from positions 1967-1976 in Figure 1H),	(Seq ID no 21)
	ACT CCG CCC CAA CTG (selected from positions 2403-2420 in Figure 1H),	(Seq ID no 22)
	CTG TCC CTA AAC CCG (selected from positions 2457-2488 in Figure 1I),	(Seq ID no 23)
20 .	TTC GAG GTT AGA TGC (selected from positions 2457-2488 in Figure 1I),	(Seq ID no 24)
	GTC CCT AAA CCC GAT (selected from positions 2457-2488 in Figure 1I),	(Seq ID no 25)
	GGT GCA CCA GAG GTT (selected from positions 2952-2956 in Figure 1I),	(Seq ID no 26)
	CTG GCG GGA CAA CTG (selected from positions 2966-2969 in Figure 1J),	(Seq ID no 27)
	TTA TCC TGA CCG AAC (selected from positions 3000-3003 in Figure 1J),	(Seq ID no 28)
25	GAC CTA TTG AAC CCG (selected from positions 3097-3106 in Figure 1J),	(Seq ID no 29)
	MTC group (16S)	
	GAA GAG ACC TTT CCG (selected from positions 76-79 in Figure 2A),	(Seq ID no 30)
	CAC TCG AGT ATC TCC (selected from positions 98-101 in Figure 2A),	(Seq ID no 31)
30	ATC ACC CAC GTG TTA (selected from positions 136-136 in Figure 2A),	(Seq ID no 32)
	GCA TCC CGT GGT CCT (selected from positions 194-201 in Figure 2B),	(Seq ID no 33)
	CAC AAG ACA TGC ATC (selected from positions 194-201 in Figure 2B).	(Seq ID no 34)
	TAA AGC GCT TTC CAC (selected from positions 222-229 in Figure 2B),	(Seq ID no 35)
	GCT CAT CCC ACA CCG (selected from position 242 in Figure 2B),	(Seq ID no 36)
35	CCG AGA GAA CCC GGA (selected from position 474 in Figure 2C),	(Seq ID no 37)
	AGT CCC CAC CAT TAC (selected from positions 1136-1145 in Figure 2C),	(Seq ID no 38)
	AAC CTC GCG GCA TCG (selected from positions 1271-1272 in Figure 2C),	(Seq ID no 39)
	GGC TTT TAA GGA TTC (selected from positions 1287-1292 in Figure 2D),	(Seq ID no 40)
	GAC CCC GAT CCG AAC (selected from position 1313 in Figure 2D),	(Seq ID no 41)
40	CCG ACT TCA CGG GGT (selected from position 1334 in Figure 2D),	(Seq ID no 42)
	, , , , , , , , , , , , , , , , , , , ,	(4·-···

MTC group (5S)

	CGG AGG GGC AGT ATC (selected from positions 86-90 in Figure 3),	(Seq ID no 43)
	Mycobacteria other than those of the MTC group (23S)	
	GAT CAA TGC TCG GTT (selected from positions 99-101 in Figure 4A),	(Seg ID no 44)
5	TTC CCC GCG TTA CCT (selected from position 183 in Figure 4A),	(Seq ID no 45)
	TTA GCC TGT TCC GGT (selected from positions 261-271 in Figure 4A),	(Seq ID no 46)
	GCA TGC GGT TTA GCC (selected from positions 281-284 in Figure 4B),	(Seq ID no 47)
	TAC CCG GTT GTC CAT (selected from positions 290-293 in Figure 4B),	(Seq ID no 48)
	GTA GAG CTG AGA CAT (selected from positions 327-335 and	,
10	343-357 in Figure 4B),	(Seq ID no 49)
	GCC GTC CCA GGC CAC (selected from positions 400-405 in	
	Figure 4B and Figure 4C),	(Seq ID no 50)
	CTC GGG TGT TGA TAT (selected from positions 453-462 in Figure 4C),	(Seq ID no 51)
	ACT ATT TCA CTC CCT (selected from positions 587-599 in Figure 4C),	(Seq ID no 52)
15	ACG CCA TCA CCC CAC (selected from positions 637-660 in Figure 4D),	(Seq ID no 53)
	CGA CGT GTC CCT GAC (selected from positions 704-712 in Figure 4D),	(Seq ID no 54)
	ACT ACA CCC CAA AGG (selected from positions 763-789 in Figure 4E),	(Seq ID no 55)
	CAC GCT TTT ACA CCA (selected from positions 1060-1074 in Figure 4E),	(Seq ID no 56)
	GCG ACT ACA CAT CCT (selected from positions 1177-1185 in Figure 4E),	(Seq ID no 57)
20	CGG CGC ATA ATC ACT (selected from positions 1259-1265 in Figure 4E),	(Seq ID no 58)
	CCA CAT CCA CCG TAA (selected from positions 1311-1327 in Figure 4F),	(Seq ID no 59)
	CGC TGA ATG GGG GAC (selected from positions 1345-1348 in Figure 4F),	(Seq ID no 60)
	GGA GCT TCG CTG AAT (selected from positions 1361-1364 in Figure 4G),	(Seq ID no 61)
	CGG TCA CCC GGA GCT (selected from positions 1361-1364 in Figure 4G),	(Seq ID no 62)
25	GGA CGC CCA TAC ACG (selected from positions 1556-1570 in Figure 4G),	(Seq ID no 63)
	GAA GGG GAA TGG TCG (selected from positions 1608-1613 in Figure 4H),	(Seq ID no 64)
	AAT CGC CAC GCC CCC (selected from positions 1626-1638 in Figure 4H),	(Seq ID no 65)
	CAG CGA AGG TCC CAC (selected from positions 1651-1659 in Figure 4H),	(Seg ID no 66)
	GTC ACC CCA TTG CTT (selected from positions 1675-1677 in Figure 4H),	(Seq ID no 67)
30	ATC GCT CTC TAC GGG (selected from positions 1734-1741 in Figure 4H),	(Seq ID no 68)
	GTG TAT GTG CTC GCT (selected from positions 1847-1853 in Figure 4I),	(Seq ID no 69)
	ACG GTA TTC CGG GCC (selected from positions 1967-1976 in Figure 4I),	(Seq ID no 70)
	GGC CGA ATC CCG CTC (selected from positions 2006-2010 in Figure 4I),	(Seq ID no 71)
	AAA CAG TCG CTA CCC (selected from positions 2025-2027 in Figure 4I),	(Seq ID no 72)
35	CCT TAC GGG TTA ACG (selected from positions 2131-2132 in Figure 4J),	(Seq ID no 73)
	GAG ACA GTT GGG AAG (selected from positions 2252-2255 in Figure 4J),	(Seq ID no 74)
	TGG CGT CTG TGC TTC (selected from positions 2396-2405 in	
	Figure 4J and Figure 4K),	(Seq ID no 75)
40	CGA CTC CAC ACA AAC (selected from positions 2416-2420 in Figure 4K),	(Seq ID no 76)
40	GAT AAG GGT TCG ACG (selected from positions 2474-2478 in Figure 4K),	(Seq ID no 77)
	ATC CGT TGA GTG ACA (selected from position 2687 in Figure 4K),	(Seq ID no 78)
	CAG CCC GTT ATC CCC (selected from position 2719 in Figure 4K),	(Seq ID no 79)

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	AAC CTT TGG GAC CTG (selected from position 2	809 in Figure 4L),	(Seq ID no 80)
	TAA AAG GGT GAG AAA (selected from positions	3062-3068 in Figure 4L).	(Seq ID no 81)
	GTC TGG CCT ATC AAT (selected from positions 3	1097-3106 in Figure 4L),	(Seq ID no 82)
5	Mycobacteria other than those of the MTC grou	up (16S)	
	AGA TTG CCC ACG TGT (selected from positions	135-136 in Figure 5A),	(Seq ID no 83)
	AAT CCG AGA AAA CCC (selected from positions	172-475 in Figure 5A),	(Seq ID no 84)
	GCA TTA CCC GCT GGC (selected from positions	1136-1144 in Figure 5A),	(Seq ID no 85)
	TTA AAA GGA TTC GCT (selected from positions 1	287-1292 in Figure 5B),	(Seq ID no 86)
10	AGA CCC CAA TCC GAA (selected from position 13	313 in Figure 5B),	(Seq ID no 87)
	GAC TCC GAC TTC ATG (selected from position 13	334 in Figure 5B),	(Seq ID no 88)
	Drug resistance		
	23S-mediated macrolide resistance (M. avium)		
15	GTC TTT TCG TCC TGC (wild-type) (selected from	positions 2568-2569	
	in Figure 6),		(Seq ID no 89)
	GTC TTA TCG TCC TGC (selected from positions 2	568 in Figure 6),	(Seq ID no 90)
	GTC TTC TCG TCC TGC (selected from positions 2	568 in Figure 6),	(Seq ID no 91)
	GTC TTG TCG TCC TGC (selected from positions 2	568 in Figure 6),	(Seq ID no 92)
20	GTC TAT TCG TCC TGC (selected from positions 2	568 in Figure 6),	(Seq ID no 93)
	GTC TCT TCG TCC TGC (selected from positions 2	568 in Figure 6),	(Seq ID no 94)
	GTC TGT TCG TCC TGC (selected from positions 2	568 in Figure 6),	(Seq ID no 95)
	16S-mediated streptomycin resistance (M. tube	rculosis)	
25	TTG GCC GGT GCT TCT (wild-type) (selected from	positions 452 in Figure 7).	(Seq ID no 96)
	TTG GCC GGT ACT TCT (selected from positions 4	52 in Figure 7),	(Seq ID no 97)
	TTG GCC GGT CCT TCT (selected from positions 4	52 in Figure 7),	(Seq ID no 98)
	TTG GCC GGT TCT TCT (selected from positions 4	52 in Figure 7),	(Seq ID no 99)
	ACC GCG GCT GCT GGC (wild-type) (selected from	positions 473-477	
30	in Figure 7),		(Seq ID no 100)
	ACC GCG GCT ACT GGC (selected from positions	473 in Figure 7),	(Seq ID no 101)
	ACC GCG GCT CCT GGC (selected from positions	473 in Figure 7), or	(Seq ID no 102)
	ACC GCG GCT TCT GGC (selected from positions	173 in Figure 7),	(Seq ID no 103)
	CGG CAG CTG GCA CGT (selected from positions	474 in Figure 7),	(Seq ID no 104)
35	CGG CCG CTG GCA CGT (selected from positions	474 in Figure 7),	(Seq ID no 105)
	CGG CTG CTG GCA CGT (selected from positions	474 in Figure 7),	(Seq ID no 106)
	CGT ATT ACC GCA GCT (selected from positions	177 in Figure 7),	(Seq ID no 107)
	CGT ATT ACC GCC GCT (selected from positions	177 in Figure 7),	(Seq ID no 108)
	CGT ATT ACC GCT GCT (selected from positions 4	177 in Figure 7),	(Seq ID no 109)
40	TTC CTT TGA GTT TTA (wild-type) (selected from p	ositions 865-866 in Figure 7),	(Seq ID no 110)
	TTC CTT TAA GTT TTA (selected from positions 86	5 in Figure 7),	(Seq ID no 111)
	TTC CTT TCA GTT TTA (selected from positions 86	5 in Figure 7),	(Seq ID no 112)

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TTC CTT TTA GTT TTA (selected from positions 865 in Figure 7),	(Seq ID no 113)
TTC CTT AGA GTT TTA (selected from positions 866 in Figure 7),	(Seq ID no 114)
TTC CTT CGA GTT TTA (selected from positions 866 in Figure 7),	(Seq ID no 115)
TTC CTT GGA GTT TTA (selected from positions 866 in Figure 7),	(Seq ID no 116)

Other examples of suitable Q subsequences are given below.

CAT GTG TCC TGT GGT and (Seq ID no 117)
CGT CAG CCC GAG AAA (Seq ID no 118)

selected so as to be complementary to M. gordonae 16S rRNA (positions 174-188 and 452-466, respectively, of GenBank entry GB:MSGRR16SI, accession no. M29563). These positions correspond to positions 192-206 and 473-487, respectively, of the alignments shown in Figure 2 and 5. Probes having this or a similar nucleobase sequence are suitable for detecting M. gordonae.

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CAC TAC ACA CGC TCG, and (Seq ID no 119)
TGG CGT TGA GGT TTC (Seq ID no 120)
selected so as to be complementary to positions 781-795 and 2369-2383, respectively, of M. kansasii 23S rRNA (GenBank entry MK23SRRNA accession number Z17212). These positions correspond to positions 774-794 and 2398-2412, respectively, of the alignments shown in Figure 1 and 4. Probes having this or a similar nucleobase sequence are suitable for detecting M. kansasii.

Precursor rRNA

25 AAC ACT CCC TTT GGA

(Seq ID no 123)

A peptide nucleic acid probe having the above-indicated nucleobase sequence is directed to M. tuberculosis precursor rRNA. The probe is complementary to positions 602 to 616 of GenBank accession number X58890.

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Especially, probes based on those nucleobase sequences with sequence identification numbers Seq ID no 62, 79 and 80 (and other probes selected from positions 1361-1364 in Figure 1F, 2719 in Figure 4K and 2809 in Figure 4L) are suitable for detecting M. avium. Probes based on the nucleobase sequence with sequence identification number Seq ID no 55 (and other probes selected from positions 763-789 in Figure 4E) are suitable for detecting M. avium, M. intracellulare and M. scrofulaceum as a group (the organisms termed the MAIS group of mycobacteria). In addition, probes based on the nucleobase sequences with sequence identification numbers Seq ID no 77 and 81 are suitable for detecting M. avium, M. intracellulare and M. paratuberculosis as a group.

The invention is further illustrated by the non-limiting examples given below.

EXAMPLES

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EXAMPLE 1

Mycobacterium species (M. bovis and M. intracellulare) 23S rDNA were partly amplified by PCR, and the PCR products were sequenced (both strands) using Cy5-labelled oligonucleotide primers (DNA Technology, Aarhus, Denmark) and the 7-deaza-dGTP Thermo Sequenase cycle sequencing kit from Amersham, Little Chalfont, England. Sequences were read using an ALFexpress automated sequencer and ALFwin (version 1.10) software from Pharmacia Biotech, Uppsala, Sweden. M. bovis and M. intracellulare 23S rRNA sequences are included at the following positions of the 23S rDNA sequence alignments: positions 681-729 (Figures 1C and 4D), positions 761-800 (Figures 1D and 4E), positions 2401-2440 (Figures 1H and 4K), positions 2441-2480 (Figures 1I and 4K), positions 2481-2520 (Figure 1I), positions 3041-3080 (Figure 4L), and positions 3081-3120 (Figures 1J and 4L).

EXAMPLE 2

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Sequence alignments (see Figures 1 to 5) of 23S, 16S and 5S rDNA of mycobacteria of the MTC group, and 23S and 16S rDNA of mycobacteria other than those of the MTC group (MOTT) were done using the Megalign (version 3.12) alignment tool from DNASTAR (Madison, WI, USA). Up to one hundred sequences were aligned at a time.

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Peptide nucleic acid probes in which the nucleobase sequence was complementary to distinctive mycobacterial rRNA were designed with due regard to secondary structures using the PrimerSelect program (version 3.04) from DNASTAR. As a control of sequence specificity, all probe sequences were subsequently matched with the GenBank and EMBL databases using BLAST sequence similarity searching at the National Center for Biotechnology Information (http://www.ncbi.nlm.nih.gov).

As examples, the following sequences were selected:

35 MTC 23S

TCA CCA CCC TCC TCC
CCA CCC TCC TCC
ACT ATT CAC ACG CGC
CCA CAC CCA CCA CAA

(Seq ID no 6) (modified Seq ID no 6) (Seq ID no 8) (Seq ID no 12)

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			•
	AAC TCC ACA CCC CCG		(Seq ID no 16)
	ACT CCA CAC CCC CGA		(Seq ID no 17)
	ACT CCG CCC CAA CTG		(Seq ID no 22)
	CTG TCC CTA AAC CCG		(Seq ID no 23)
5	TTC GAG GTT AGA TGC		(Seq ID no 24)
	GTC CCT AAA CCC GAT		(Seq ID no 25)
	GAC CTA TTG AAC CCG		(Seq ID no 29)
	MTC 16S		
10	GCA TCC CGT GGT CCT		(Seq ID no 33)
	CAC AAG ACA TGC ATC		(Seq ID no 34)
	GGC TTT TAA GGA TTC		(Seq ID no 40)
	MOTT 23S		
15	GAT CAA TGC TCG GTT		(Seq ID no 44)
	CGA CTC CAC ACA AAC		(Seq ID no 76)
	MOTT 16S		
	GCA TTA CCC GCT GGC		(Seq ID no 85)
20			(004 15 110 00)
	Drug resistance		
	GTC TTA TCG TCC TGC		(Seq ID no 90)
	GTC TTC TCG TCC TGC		(Seq ID no 91)
	GTC TTG TCG TCC TGC		(Seq ID no 92)
25	GTC TAT TCG TCC TGC		(Seq ID no 93)
	GTC TCT TCG TCC TGC		(Seq ID no 94)
	GTC TGT TCG TCC TGC		(Seq ID no 95)
	Precursor rRNA		
30	AAC ACT CCC TTT GGA		(Seq ID no 123)
	Non-sense probes		
	GTC CGT GAA CCC GAT		/D IP
	TAC GCT CTT TGA GCT		(Seq ID no 121)
35			(Seq ID no 122)
	EXAMPLE 3		

Peptide nucleic acid probes were synthesised using an Expedite 8909 Nucleic Acid Synthesis System purchased from PerSeptive Biosystems (Framingham, USA). The peptide nucleic acid probes were terminated with two $\beta\mbox{-alanine}$ molecules or with one or two lysine molecule(s) and, before cleavage from the resin, labelled with 5-(or 6)-carboxyfluorescein (Flu) or

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(OK 745/modified Seq ID no 89)

(OK 746/modified Seq ID no 90)

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rhodamine (Rho) at the $\beta\text{-amino}$ group of alanine (peptide label) or $\epsilon\text{-amino}$ group of lysine (peptide label), respectively. Probes were purified using reverse phase HPLC at 50°C and characterised using a G2025 A MALDI-TOF MS instrument (Hewlett Packard, San Fernando, California, USA). Molecular weights determined were within 0.1% of the calculated molecular weights.

The following labelled peptide nucleic acid probes were synthesised:

Drug resistance

Lys(Flu)-GTC TTT TCG TCC TGC-NH2

Lys(Rho)-GTC TTA TCG TCC TGC-NH₂

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	MTC 23S	
10	Lys(Flu)-Lys(Flu)-TCA CCA CCC TCC TCC-NH2	(OK 446/modified Seq ID no 6)
	Lys(Flu)-Lys(Flu)-CCA CCC TCC TCC-NH ₂	(OK 575/modified Seq ID no 6)
	Lys(Flu)-Lys(Flu)-ACT ATT CAC ACG CGC-NH2	(OK 447/modified Seq ID no 8)
	Lys(Flu)-ACT ATT CAC ACG CGC-NH ₂	(OK 688/modified Seq ID no 8)
	Lys(Flu)-Lys(Flu)-CCA CAC CCA CCA CAA-NH₂	(OK 448/modified Seq ID no 12)
15	Lys(Flu)-Lys(Flu)-AAC TCC ACA CCC CCG-NH ₂	(OK 449/modified Seq ID no 16)
	Lys(Flu)-Lys(Flu)-ACT CCA CAC CCC CGA-NH ₂	(OK 309/modified Seq ID no 17)
	Lys(Flu)-Lys(Flu)-ACT CCG CCC CAA CTG-NH₂	(OK 450/modified Seq ID no 22)
	Lys(Flu)-Lys(Flu)-CTG TCC CTA AAC CCG-NH ₂	(OK 305/modified Seq ID no 23)
	Lys(Flu)-Lys(Flu)-TTC GAG GTT AGA TGC-NH₂	(OK 306/modified Seq ID no 24)
20	Lys(Flu)-TTC GAG GTT AGA TGC-NH ₂	(OK 682/modified Seq ID no 24)
	Lys(Flu)-Lys(Flu)-GTC CCT AAA CCC GAT-NH ₂	(OK 307/modified Seq ID no 25)
	Lys(Flu)-GTC CCT AAA CCC GAT-NH2	(OK 654/modified Seq ID no 25)
	Lys(Flu)-GAC CTA TTG AAC CCG-NH₂	(OK 660/modified Seq ID no 29)
25	MTC 16S	
	Lys(Flu)-Lys(Flu)-Gly-GCA TCC CGT GGT CCT-NH2	(OK 223/modified Seq ID no 33)
	Lys(Flu)-Lys(Flu)-CAC AAG ACA TGC ATC-NH₂	(OK 310/modified Seq ID no 34)
	Lys(Flu)-CAC AAG ACA TGC ATC-NH2	(OK 655/modified Seq ID no 34)
	Lys(Flu)-GGC TTT TAA GGA TTC-NH₂	(OK 689/modified Seq ID no 40)
30	Lys(Rho)-GGC TTT TAA GGA TTC-NH2	(OK 702/modified Seq ID no 40)
	MOTT 23S	,
	Flu-β-Ala-β-Ala-GAT CAA TGC TCG GTT-NH ₂	(OK 624/modified Seq ID no 44)
	Flu-β-Ala-β-Ala-CGA CTC CAC ACA AAC-NH ₂	(OK 612/modified Seq ID no 76)
35		,
	MOTT 16S	
	Flu-β-Ala-β-Ala-GCA TTA CCC GCT GGC-NH₂	(OK 623/modified Seq ID no 85)

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Lys(Rho)-GTC TTC TCG TCC TGC-NH₂	(OK 746/modified Seq ID no 91)
Lys(Rho)-GTC TTG TCG TCC TGC-NH₂	(OK 746/modified Seq ID no 92)
Lys(Rho)-GTC TAT TCG TCC TGC-NH₂	(OK 747/modified Seq ID no 93)
Lys(Rho)-GTC TCT TCG TCC TGC-NH ₂	(OK 747/modified Seq ID no 94)
Lys(Rho)-GTC TGT TCG TCC TGC-NH ₂	(OK 747/modified Seq ID no 95)

Precursor rRNA

Lys(Fiu)-AAC ACT CCC TTT GGA-NH2

(OK 749/modified Seg ID no 123)

10 Reduction of non-specific binding

GTC CGT GAA CCC GAT-NH₂
Gly-TAC GCT CTT TGA GCT-NH₂

(OK 507/modified Seq ID no 121) (OK 714/modified Seq ID no 122)

EXAMPLE 4

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Initially the ability of the peptide nucleic acid probes to react with target sequences of mycobacterial rRNA was tested by dot blot carried out with rRNA from M. bovis BCG, M. avium and E.coli.

M. bovis BCG (Statens Serum Institut, Denmark) and M. intracellulare (kindly provided by Statens Serum Institut) were grown in Dubos broth (Statens Serum Institut) or on Löwenstein-Jensen slants (Statens Serum Institut) at 37 °C. RNA was isolated from the bacterial cells using TRI-reagent (Sigma) following manufacture's directions. E. coli rRNA was purchased from Boehringer Mannheim, Germany.

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200 ng M. bovis RNA, M. intracellulare RNA and E. coli rRNA were dotted onto membranes (Schleicher & Schüel, NY 13 N), and the membranes were dried and fixed under UV light for 2 minutes.

30 Protocol for dot blot assay

Each of the probes (70 nM probe in hybridisation solution (50 mM Tris, 10 mM NaCl, 10% (w/v) Dextran sulphate, 50% (v/v) glycerol, 5 mM EDTA, 0.1% (w/v) sodium pyrophosphate, 0.2% (w/v) polyvinylpyrrolidone, 0.2% (w/v) Ficoll, pH 7.6.)) were spotted onto a membrane. Hybridisation was continued for 1.5 hours at 55 or 65 °C, respectively. The membranes were rinsed 2 times for 15 minutes in 2 × SSPE buffer (1 x SSPE: 0.15 M NaCl, 10 mM sodium phosphate, 1 mM EDTA, pH 7.4) containing 0.1% SDS at ambient temperature, and subsequently 2 times for 15 minutes in 0.1 × SSPE buffer containing 0.1% SDS at 55 or 65 °C (see Table 1). The membrane was blocked with 0.5% (w/v) casein dissolved in 0.5M NaCl, 0.05M Tris/HCl pH 9.0. Thereafter, the membranes were incubated for 1 hour with rabbit-anti

FITC antibody labelled with alkaline phosphatase (AP) (DAKO K0046 vial A) diluted 1:2000 in 0.5% casein dissolved in 0.5M NaCl, 0.05M Tris/HCl pH 9.0. After incubation, the membranes were washed 3 times 5 minutes with TST buffer (0.05M Tris, 0.5M NaCl, 0.5% (w/v) Tween 20[®], pH 9) at ambient temperature. Bound probes were visualised following standard procedures using BClP/NBT, and the visualisation was stopped by incubation for 10 minutes with 10 mM EDTA. The blot was dried at 50 °C.

The results are given in Table 1 below.

TABLE 1

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	E. coli rRNA		M. bovis BCG RNA			cellulare NA
Probe	55 °C	65 °C	55 °C	65 °C	55 °C	65 °C
OK 305	negative	negative	positive	positive	negative	weak
OK 307	negative	negative	positive	positive	negative	weak
OK 309	negative	negative	positive	positive	negative	weak
OK 223	negative	negative	positive	positive	nd	nd
OK 310	negative	negative	negative	positive	negative	negative

nd: Not determined

The results indicate that all five peptide nucleic acid probes are capable of hybridising to target sequence of M. bovis BCG rRNA (as a representative of the MTC group), whereas no hybridisation to E. coli rRNA (as a representative of organisms other than mycobacteria) and no detectable hybridisation to M. intracellulare rRNA were observed (as a representative of the MOTT group).

EXAMPLE 5

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This example illustrates the ability of the peptide nucleic acid probes to penetrate the mycobacterial cell wall and subsequently hybridise to target sequence of mycobacteria of the MTC group and not mycobacteria of the MOTT group, in particular not mycobacteria of the MAC group, or Neisseria gonorrhoeae, by fluorescence *in situ* hybridisation (FISH).

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Preparation of bacterial slides

M. bovis BCG (Statens Seruminstitut, Denmark), M. avium (kindly provided by Statens Seruminstitut, Denmark), and M. intracellulare (kindly provided by Statens Seruminstitut,

Denmark) were grown in Dubos broth (Statens Seruminstitut, Denmark) or on Löwenstein-Jensen slants (Statens Seruminstitut, Denmark) at 37 °C. N. gonorrhoeae (Statens Seruminstitut, Denmark) was grown on chocolate agar (Statens Seruminstitut, Denmark) at 37 °C with additional 5% CO₂.

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Cultures were smeared onto microscope slides and fixed according to standard procedures. Prior to the hybridisation, the smears were immersed into 80% ethanol for 15 minutes, and subsequently rinsed with water and air dried. This step is not essential for the following hybridisation step, but it is anticipated that it will kill any viable mycobacteria on the slides, and may further serve as an additional fixation step.

Protocol for fluorescence in situ hybridisation (FISH)

- The bacterial slide was covered with a hybridisation solution containing the probe in question.
- 15 2. The slide was incubated in a humid incubation chamber at 45°C or 55°C for 90 minutes.
 - The slide was washed 25 minutes at 45°C or 55°C in prewarmed wash solution (5 mM
 Tris, 145 mM NaCl, pH 10) followed by 30 seconds in water.
 - 4. The slide was dried and mounted with IMAGEN Mounting Fluid (DAKO, Copenhagen, Denmark)

The hybridisation solution contains 50 mM Tris, 10 mM NaCl, 10% (w/v) Dextran sulphate, 30% (v/v) formamide, 0.1% (v/v) Triton X-100[®], 5 mM EDTA, 0.1% (w/v) sodium pyrophosphate, 0.2% (w/v) polyvinylpyrrolidone, 0.2% (w/v) Ficoll, pH 7.6.

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Whenever possible, the applied equipment was heat-treated, and solutions were exposed to 1µI/ml diethylpyrocarbonate (Sigma Chemical Co.) in order to inactivate nucleases.

Microscopically examinations were conducted using a fluorescence microscope (Leica, Wetzlar, Germany) equipped with a 100×/1.20 water objective, a HBO 100 W lamp and a FITC filter set. Mycobacteria were identified as fluorescent, 1 - 10 μm slender, rod-shaped bacilli.

Fluorescein-labelled peptide nucleic acid probes targeting 23S rRNA of the mycobacteria of the MTC group (OK 306, OK 309, OK 446, OK 449) and 16S rRNA of the mycobacteria of the MTC group (OK 310) were tested. Individual probe concentrations and incubation temperatures are listed together with the results in Table 2 and 3.

TABLE 2

	OK 306	OK 309	OK 446	OK 449
	250nM	250nM	500nM	500nM
	45°C	45°C	55°C	55°C
M. bovis BCG	positive	positive	positive	positive
M. avium	negative	negative	negative	negative
M. intracellulare	negative	negative	not determined	not determined
N. gonorrhoeae	negative	negative	not determined	not determined

TABLE 3

	OK 447	OK 310	OK 306/OK 310
	1μМ	250nM	500/500nM
	55°C	45°C	55°C
M. bovis BCG	positive	positive	positive
M. avium	negative	negative	negative
M. intracellulare	not determined	negative	negative
N. gonorrhoeae	not determined	negative	not determined

5 It can be concluded that the probes are able to penetrate the mycobacterial cell wall of mycobacterium cultures and subsequently hybridise to target rRNA sequence. This makes possible the development of fluorescence in situ hybridisation (FISH) protocols for specific detection of mycobacteria.

10 EXAMPLE 6

Test of probes on clinical smears of sputum

The ability of the peptide nucleic acid to penetrate the cell wall of mycobacteria of the MTC group in clinical samples was tested on smears of sputum from suspected cases of tuberculosis (kindly provided by Division of Microbiology, Ramathibodi Hospital, Bangkok, Thailand) by fluorescence in situ hybridisation (FISH). Smears from the same patient were initially evaluated positive by Ziehl-Neelsen staining, which shows only the presence of acid fast bacilli, not whether these are mycobacteria of the MTC group.

Fluorescein-labelled peptide nucleic acid probes targeting 23S rRNA of the mycobacteria of the MTC group (OK 306, OK 446, OK 449) and 16S rRNA of the mycobacteria of the MTC group (OK 310) were used. Furthermore, a random peptide nucleic acid probe (a 15-mer wherein each position may be A, T, C or G (obtained from Millipore Corporation, Bedford, MA, USA) was added to the hybridisation solution in order to increase the signal-to-noise ratio.

FISH was carried out at 55 °C as described in Example 5. Applied probe concentrations are listed together with the results in Table 4 and 5.

TABLE 4

Sample	OK 446/Random	OK 449/Random	Ziehl-Neelsen
number	1μΜ/50μΜ	1μΜ/50μΜ	staining
285	Positive	Positive	4+
335	Positive	Eq.	2+
345	Positive	Positive	3+
224	Positive	Positive	3+
297	Negative	Eq.	2+
179	Negative	Negative	4+
247	Negative	Negative	2+
255	Positive	Positive	2+
202	Eq.	Positive	2+

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TABLE 5

Sample	OK 306/OK 310	Ziehl-Neelsen
number	500/500 nM	staining
213 .	Positive	4+
292	Positive	4+
159	Positive	3+
287	Positive	3+

Smears stained by Ziehl-Neelsen staining were examined with a 100x objective and scored according to the following method: -: 0 bacilli, +/-: 1-200 per 300 fields, 2+: 1-9 per 10 fields, 3+: 1-9 per field, 4+: >9 per field.

Positive: Several mycobacteria were identified in the smear. Negative: No fluorescent mycobacteria were identified in the smear. Eq. Few (1-3) fluorescent mycobacteria were identified in the smear.

It appears from the table that the peptide nucleic acid probes are able to penetrate and subsequently hybridise to target sequence of mycobacteria of the MTC-group in AFB-positive sputum smears. The fact that not all AFB-positive sputum smears are found positive with applied probes indicate that not all AFB-positive sputum smears contains mycobacteria of the MTC-group.

EXAMPLE 7

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The reactivity and specificity of selected peptide nucleic acid probes for detecting

mycobacteria of the MTC group as well as probes for detecting mycobacteria of the MOTT group were evaluated by fluorescence in situ hybridisation (FISH) on control smears prepared from cultures of different mycobacterium species. The mycobacterium species were selected so as to be representative for the mycobacterium genus as well as to include clinically relevant species.

M. tuberculosis (ATCC 25177), M. bovis BCG (ATCC 35734), M. intracellulare (ATCC 13950), M. avium (ATCC 25292), M. kansasii (ATCC12479), M. gordonae (ATCC 14470), M. scrofulaceum (ATCC 19981), M. abscessus (ATCC19977), M. marinum (ATCC 927), M. simiae (ATCC 25575), M. szulgai (ATCC 35799), M. flavescens (ATCC 23033), M. fortuitum (ATCC 43266) and M. xenopi (ATCC19250) were grown at Dubos broth (Statens Serum Institut) at 37 °C with the exception of M. marinum which was grown at 32 °C.

Smears were prepared as described in Example 5. FISH was carried out as described below.

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Protocol for fluorescence in situ hybridisation (FISH)

- The bacterial slide was covered with a hybridisation solution containing the probe in question.
- 2. The slide was incubated in a humid incubation chamber at 55°C for 90 minutes.
- The slide was washed 30 minutes at 55°C in prewarmed wash solution (5 mM Tris, 15 mM NaCl, 0.1% (v/v), Triton X-100[®], pH 10) followed by 30 seconds in water.
 - The slide was dried and mounted with IMAGEN Mounting Fluid (DAKO, Copenhagen, Denmark)
- The hybridisation solution contained 50 mM Tris, 10 mM NaCl, 10% (w/v) Dextran sulphate, 30% (v/v) formamide, 0.1% (v/v) Triton X-100°, 5 mM EDTA, 0.1% (w/v) sodium pyrophosphate, 0.2% (w/v) polyvinylpyrrolidone, and 0.2% (w/v) Ficoll, pH 7.6. To avoid non-specific binding of the labelled peptide nucleic acid probe, 1-5 μM of non-labelled, non-sense peptide nucleic acid probe was added to the hybridisation solution (OK 507/modified Seq ID no 121 and/or OK 714/modified Seq ID no 122).

Whenever possible, the applied equipment was heat-treated, and solutions were exposed to 1µl/ml diethylpyrocarbonate (Sigma Chemical Co.) in order to inactivate nucleases.

Microscopic examinations were conducted using a fluorescence microscope (Leica, Wetzlar, Germany) equipped with a 100×/1.30 oil objective, a HBO 100 W lamp and a FITC/TRITC dual band filter set. Mycobacteria were identified on basis of both fluorescence (strong, medium, weak, no) and morphology (1-10 μm slender, rod-shaped bacilli. Mycobacteria of the MOTT

group may appear pleomorphic, ranging in appearance from long rods to coccoid forms)

Probe concentrations are listed together with the results in Table 6 and 7 (probes targeting mycobacteria of the MTC group) and Table 8 (probes targeting to mycobacteria of the MOTT group).

TABLE 6

	OK 450	OK 682	OK 689	OK 688	OK 660
	25 nM	100 nM	100 nM	250 nM	100 nM
M. tuberculosis	+++	+++	+++	+++	+++
M. bovis BCG	+++	+++	+++	+++	+++
M. intracellulare		-	-	-	
M. avium	-	•	-	-	•
M. kansasii	++	-	-	-	-
M. gordonae	-	-	-	-	-
M. scrofulaceum	+++	-	-	-	•
M. abscessus	_	-	-	-	+
M. marinum	+++	•	+	+	+++
M. simiae	-	•	-	-	-
M. szulgai	+++	-	-	-	-
M. flavescens	-	++	-	-	-
M. fortuitum		+		-	-
M. xenopi	-	++			

⁺⁺⁺ strong fluorescence, ++ medium fluorescence, + weak fluorescence, - no fluorescence

TABLE 7

Mycobacteria	OK 655	OK 448	OK 654	OK 446
	150 nM	50 nM	100 กฟ้	25 nM
M. tuberculosis	+++	+++	+++	+++
M. bovis BCG	+++	+++	+++	+++
M. intracellulare	-	-	•	•
M. avium	-	-	-	•
M. kansasii	-	•	-	-
M. gordonae	-	-	•	-
M. scrofulaceum	-	-	-	-
M. abscessus		-	+	•
M. marinum	-	•	+	+++
M. simiae	-	. •	-	•
M. szulgai	1 - 1	-	•	-
M. flavescens	-	-	-	•
M. fortuitum	-		•	•
M. xenopi	-		-	-

⁺⁺⁺ strong fluorescence, ++ medium fluorescence, + weak fluorescence, - no fluorescence

TABLE 8

Mycobacteria	OK 612	OK 624	OK 623
in y cobactoria	OKO12 .	OK 024	UK 023
	100 nM	100 nM	100 nM
M. tuberculosis	-	-	-
M. bovis BCG		-	-
M. intracellulare	-	++	++
M. avium	+++	+++	+++
M. kansasii	-	-	+++
M. gordonae	-	++	++ ·
M. scrofulaceum	-	++	++
M. abscessus	-	++	+++
M. marinum	-	•	-
M. simiae	•	++	+++
M. szulgai	-	-	+++
M. flavescens		•	-
M. fortuitum	-	++	-
M. xenopi	-	-	•

⁺⁺⁺ strong fluorescence, ++ medium fluorescence, + weak fluorescence, - no fluorescence

Each of probes indicated in Table 6, 7 and 8 was further investigated with regard to hybridisation to other common respiratory bacteria, namely Corynebacterium spp.,

Fusobacterium nucleatum, Haemophilus influenzae, Klebsiella pneumoniae. Pseudomonas aeruginosa, Propionibacterium acnes, Streptococcuc pneumoniae, Staphylococcus aureus, Brahamella catarrahalis, Escherichia coli, Neisseria spp., Actinobacter calcoaceticus, Actinomyces spp., Enterobacter aerogenes, Proteus mirabilis, Pseudomonas maltophilia, Streptocussuc viridans, and Norcardia asteroides. No cross-hybridisation was observed by fluorescence in situ hybridisation to any of these bacteria in the case of OK 682, OK 654, OK 655, OK 688, OK 660, OK 612, OK 624 and OK 623. Some cross-reactivity was observed in the case of OK 446 (to P. acnes), OK 448 (to P. acnes and B. catarrhalis), and OK 450 (to P. acnes and B. catarrhalis).

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Table 6 and 7 shows that none of the MTC probes cross-react with M. intracellulare and/or M. avium, but indeed strongly with M. tuberculosis and M. bovis BCG. As shown in Table 8, both OK 624 and OK 623 hybridise to M. intracellulare and M. avium which are both members of the MAC group, whereas none of them hybridise to M. tuberculosis or M. bovis BCG. OK 612 hybridises to M. avium only. It should be noted that the aligned sequence of M. intracellulare has just one nucleobase difference to the target sequence of M. avium, see Figure 4K.

The data support the use of the methodology described in claim 3 and 4 and exemplified in Example 2 for design of peptide nucleic acid probes that are capable of hybridising to target sequence of one or more mycobacterium species and not to other mycobacterium species having at least one nucleobase difference to the target sequence.

EXAMPLE 8

To study the usefulness of the peptide nucleic acid probes in distinguishing between mycobacteria of the MTC group and mycobacteria of the MOTT group, the probes were tested on smears of mycobacterium-positive cultures prepared from 34 + 28 clinical samples (sputum samples, other respiratory samples and extrapulmonary samples) from individuals suspected of tuberculosis or other mycobacterial infections (kindly provided by the Mycobacterium Department, Statens Serum Institut, Denmark). Complex/species identification data obtained with the AccuProbe tests from Gen-Probe Inc., USA were available for each sample.

Table 9 shows the results obtained with four different peptide nucleic acid probes targeting mycobacteria of the MTC group (OK 682, OK 660, OK 688 and OK 689) and one probe targeting mycobacteria of the MOTT group (OK 623), and Table 10 shows the results obtained with two peptide nucleic acid probes targeting mycobacteria of the MOTT group (OK 623 and OK 612) and a mixture of two probes targeting mycobacteria of the MTC group (OK 688 and OK 689). Data are arranged according to the results obtained by AccuProbe. Sample

preparation, hybridisation and visualisation were performed as described in Example 7.

TABLE 9

Complex/	OK 623	OK 682	OK 660	OK 688	OK 689
species (n)	25 nM	100 nM	100 nM	250 nM	100 nM
	n _p				
MTC (23)	0	23	23	23	23
M. avium (5)	5	0	0	0	0
M. gordonae (3)	3	0	0	0	0
Unknown (3)	3	0	0	0	0

n_p denotes number of positive samples.

The term "unknown" means that the sample not contains mycobacteria of the MTC group, or mycobacteria of the MAC group according the AccuProbe test, but further species identification was not performed.

TABLE 10

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Complex/	OK 623	OK 612	OK 688/OK 689	
species (n)	25nM	100 nM	50 nM/50 nM	
	n _e	n _p	n _p	
MTC (17)	0		16	
M. avium (2)	2	2	0	
M. gordonae (4)	3	0	0	
Unknown (5)	5	0	0	

n_p denotes number of positive samples.

- The term "unknown" means that the sample not contains mycobacteria of the MTC group, or mycobacteria of the MAC group according to the AccuProbe test, but further species identification was not performed.
- The results shown in Table 9 are in conformity with the complex/species identification performed with the AccuProbe tests, and thus confirm that peptide nucleic acid probes can be used to determine whether an infection is caused by mycobacteria of the MTC group or by mycobacteria of the MOTT group.
- From the results in Table 10, it can be seen that it is possible to differentiate between mycobacteria of the MTC group and mycobacteria of the MOTT group with 100% specificity and 91-94% sensitivity relative to results obtained by the AccuProbe tests. Furthermore, OK 612 is very suitable for specific identification of M. avium among those being positive for mycobacteria of the MOTT group as the result is positive in the case of M. avium and negative in the other cases of mycobacteria of the MOTT group.

EXAMPLE 9

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Direct detection of mycobacteria in clinical smears of sputum

This example demonstrates the ability of the peptide nucleic acid to detect and identify mycobacteria directly in AFB-positive sputum samples from suspected cases of tuberculosis (kindly provided by Division of Microbiology, Ramathibodi Hospital, Bangkok, Thailand) and suspected cases of other mycobacterial infections (kindly provided by Clinical Microbiology Dept., Rigshospitalet, Copenhagen, Denmark) by FISH is shown.

The clinical smears were prepared according to the procedure described in Example 5, and FISH was performed as described in Example 7. The results are shown in Table 11.

TABLE 11

	OK 623	OK 654	OK 655	OK 682	OK 688	OK 689
Sample no.	25 nM	100 nM	150 nM	100 nM	250 nM	100 nM
1	•	++	++	++	++	++
175	-	++	nd	nd	++	++
459	-	-	nd	nd	. •	-
166	-	-	-	nd	-	-
268	-	++	++	++	++	++
34267	++	-		-	-	

nd: not determined

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+++ strong fluorescence, ++ medium fluorescence, + weak fluorescence, - no fluorescence

It appears from examples in Table 11 that AFB-positive sputum smears were evaluated positive for mycobacteria of the MTC group (sample numbers 1, 175, and 268), positive for mycobacteria of the MOTT group (sample number 37267), or negative for mycobacteria (sample numbers. 459 and 166) by the applied probes. Thus, PNA-probes are useful reagents for specific identification of mycobacteria directly in sputum smears by fluorescence in situ hybridisation. AFB-positive sputum samples that are negative with all probes may be explained in three ways: a) the sample may contain mycobacteria not detected by the probes, e.g. M. fortuitum, b) the sample may contain other acid-fast bacteria than mycobacteria, or c) the mycobacteria in the sample lack or have a strongly reduced content of rRNA due to for example antibiotic treatment.

In conclusion, direct identification of mycobacteria in smear-positive sputum samples by peptide nucleic acid-based fluorescence in situ hybridisation combines simplicity and morphological advantages of current staining methods with concominant species identification, and will thus allow clinical microbiology laboratories to benefit from the

advantages offered by molecular techniques to provide crucial information pertaining to therapy and patient management.

EXAMPLE 10

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This example demonstrates simultaneous detection and identification of mycobacteria of the MTC group and mycobacteria of the MOTT group using differently labelled probes targeting mycobacteria of the MTC group and mycobacteria of the MOTT group, respectively, by fluorescence in situ hybridisation.

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Control smears of different mycobacterium species were prepared as described in Example 5. In addition, smears containing a mixture of M. tuberculosis and M. avium were prepared (Table 8, last row). FISH was performed as described in Example 7.

A rhodamine-labelled peptide nucleic acid probe targeting 16S rRNA of mycobacteria of the MTC group (OK 702) and a fluorescein-labelled peptide nucleic acid probe targeting 16S rRNA of mycobacteria of the MOTT group (OK 623) were applied simultaneously in the concentrations listed in Table 12 together with the results.

20 TABLE 12

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Mycobacterium species	OK 623/OK 702	
2.5	25/250 nM	
M. tuberculosis	- (G)/ +++ (R)	
M. bovis BCG	- (G)/ +++ (R)	
M. avium	+++ (G)/ - (R)	
M. intracellulare	+++ (G)/ - (R)	
M. kansasii	+++ (G)/ - (R)	
M. avium / M. tuberculosis	+++ (G)/+++ (R)	

⁺⁺⁺ strong fluorescence - no fluorescence

G green fluorescence, R red fluorescence

Mycobacteria of the MTC group, i.e. M. tuberculosis and M. bovis, were observed as green fluorescent mycobacteria, whereas mycobacteria of the MOTT group, i.e. M. avium, M. intracellulare and M. kansasii, were observed as red fluorescent mycobacteria. Mycobacteria in the M. avium/M. tuberculosis mixture were identified by a mixture of both green fluorescent mycobacteria and red fluorescent mycobacteria.

30 The results show that it is possible to distinguish between different Mycobacterium species in

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one smear using a mixture of differently labelled probes. Such simultaneous detection and identification of mycobacteria may further be extended to comprise three or more differently labelled peptide nucleic acid probes.

5 EXAMPLE 11

The ability of a peptide nucleic acid probes to hybridise to precursor rRNA and further to distinguish between precursor rRNA of M. tuberculosis and precursor rRNA of M. avium was investigated by fluorescence in situ hybridisation.

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Smears were prepared as described in Example 5 and FISH were carried out as described in Example 7 using a fluorescein-labelled probe targeting precursor rRNA of M. tuberculosis (OK 749). The results are given in Table 13.

15 TABLE 13

Mycobacterium	OK 749
	1000 nM
M. tuberculosis	+
M. avium	-

⁺ weak fluorescence - no fluorescence

From the results, it can be concluded that it is possible to detect precursor rRNA, and further that is possible to distinguish between precursor rRNA from different mycobacterium species. The application of peptide nucleic acid targeting precursor rRNA may be particularly useful for measuring the mycobacterial growth and thus be an indicator of the viability of the mycobacteria. This would in particular be important for monitoring of the effect of antibiotics in relation to both treatment of tuberculosis and drug susceptibility studies.

25 EXAMPLE 12

The ability of peptide nucleic acid probes for differentiation of drug susceptible and drug resistant mycobacteria was evaluated using a fluorescein-labelled probe targeting the wild type sequence of 23S rRNA of M. avium and M. intracellulare together with rhodamine-labelled probes targeting single point mutations associated with macrolide resistance in M. avium and M. intracellulare.

Smears were prepared as described in Example 5 from cultures of M. avium (ATCC no. 25292) and M. intracellulare (ATCC no. 13950). These strains are anticipated to contain the

wild type sequence of rRNA. Macrolide resistant variants were not available. FISH was carried out as described in Example 7 using a fluorescein-labelled peptide nucleic acid probe targeting wild type 23S rRNA (OK 745) and a mixture of rhodamine-labelled peptide nucleic acid probes targeting the three possible mutations at position 2568 (OK 746) and at position 2569 (OK 747) of M. avium 23S rDNA of GenBank entry X52917 (see Figure 6). The results are given in Table 14.

TABLE 14

Mycobacterium species	OK 745/OK 746/OK 747
	500/500/500 nM
M. avium (wild type)	+++ (G)/ - (R)
M. intracellulare (wild type)	+++ (G)/ - (R)

⁺⁺⁺ strong fluorescence - no fluorescence

10 G green fluorescence, R red fluorescence

OK 746 and OK 747 are each a mixture of three single point mutation probes

The results in Table 14 show that M. avium and M. intracellulare are detected with the fluorescein-labelled probe (OK 745) targeting M. avium and M. intracellulare wild types and not detected with the mixture of rhodamine-labelled probes (OK 746 and OK 747) targeting single point mutations associated with macrolide resistance. Such peptide nucleic acid probes targeting the wild type and drug resistant variants, respectively, may be important tools for both the prediction of an efficient therapy as well as for monitoring the effect of the treatment.

EXAMPLE 13

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To illustrate the speed with which peptide nucleic acid probes penetrate the mycobacterial cell wall and subsequently hybridise to their target sequence the protocol described in Example 7 was modified to 15 minutes hybridisation time and the results compared with 90 minutes hybridisation time. Smears were prepared as described in Example 5. The results are given in Table 15.

TABLE 15

	OK 623 25 nM		OK 689		
			100 nM		
	15 min	90 min	15 min	90 min	
M. tuberculosis			++	+++	
M. avium	++	+++			

⁺⁺⁺ strong fluorescence ++ medium fluorescence

The data presented in Table 15 show that hybridisation by peptide nucleic acid probes inside the mycobacterial cells is accomplished in a very short time resulting in a detectable signal after just 15 minutes incubation. Thus, the use peptide nucleic acid probes makes possible the development of very fast fluorescence in situ hybridisation protocols.

10 EXAMPLE 14

To describe the ability of very short peptide nucleic acid probes to hybridise to target sequences, a 12-mer peptide nucleic acid probe labelled with fluorescein (OK 575) was tested by fluorescence in situ hybridisation (FISH).

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Smears were prepared as described in Example 5 and FISH were carried out as described in Example 7. The results are given in Table 16.

TABLE 16

Mycobacterium	OK 575
	50 nM
M. tuberculosis	+
M. bovis BCG	++
M. avium	•
M. intracellulare	•
M. kansasii	-

20 ++ medium fluorescence + weak fluorescence - no fluorescence

The results in table 17 shows that a 12-mer peptide nucleic acid probe is capable of hybridising specifically to target sequences under the same stringency conditions as 15-mers. A lower florescence intensity is obtained as the T_m for a 12-mer peptide nucleic acid probe is lower than T_m for a 15-mer peptide nucleic acid probe.

⁺ weak fluorescence - no fluorescence

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The data clearly suggest that by lowering the stringency condition, e.g. by decreasing the hybridisation/washing temperature and/or the concentration of formamide, even shorter probes may be applied for detection of mycobacteria provided that specific sequences of such can be designed.

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CLAIMS

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- 1. Peptide nucleic acid probe for detecting a target sequence of one or more mycobacteria optionally present in a sample, said probe being capable of hybridising to a target sequence of mycobacterial rDNA, precursor rRNA or rRNA forming detectable hybrids, and a mixture of such probes.
- 2. Peptide nucleic acid probe according to claim 1, said probe being capable of hybridising to a target sequence of mycobacterial rDNA, precursor rRNA, or 23S, 16S or 5S rRNA forming detectable hybrids, and a mixture of such probes.
- 3. Peptide nucleic acid probe according to claim 1 or 2, said probe being capable of hybridising to a target sequence of mycobacterial rDNA, precursor rRNA, or 23S, 16S or 5S rRNA forming detectable hybrids, said target sequence being obtainable by
- (a) comparing the nucleobase sequences of said mycobacterial rRNA or rDNA of one or more mycobacteria to be detected with the corresponding nucleobase sequence of organism(s), in particular other mycobacteria, in particular other mycobacteria, from which said one or more mycobacteria are to be distinguished,
- (b) selecting a target sequence of said rRNA or rDNA which includes at least one nucleobase differing from the corresponding nucleobase of the organism(s), in particular other mycobacteria, from which said one or more mycobacteria are to be distinguished, and
- (c) determining the capability of said probe to hybridise to the selected target sequence to form detectable hybrids, and a mixture of such probes.
- 4. Peptide nucleic acid probe according to claim 1 or 2, said probe being capable of hybridising to a target sequence of mycobacterial rDNA, precursor rRNA or 23S, 16S or 5S rRNA forming detectable hybrids, said probe being obtainable by
- (a) comparing the nucleobase sequences of said mycobacterial rRNA or rDNA of one or more mycobacteria to be detected with the corresponding nucleobase sequence of organism(s), in particular other mycobacteria, in particular other mycobacteria, from which said one or more mycobacteria are to be distinguished,

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- (b) selecting a target sequence of said rRNA or rDNA which includes at least one nucleobase differing from the corresponding nucleobase of the organism(s), in particular other mycobacteria, from which said one or more mycobacteria are to be distinguished.
- 5 (c) synthesising said probe, and
 - (d) determining the capability of said probe to hybridise to the selected target sequence to form detectable hybrids, and a mixture of such probes.
 - 5. Peptide nucleic acid probe according to any one of claims 1 to 4 for detecting a target sequence of one or more mycobacteria of the Mycobacterium tuberculosis Complex (MTC) or for detecting a target sequence of one or more mycobacteria other than mycobacteria of the Mycobacterium tuberculosis Complex (MOTT) optionally present in a sample, which probe comprises from 6 to 30 polymerised peptide nucleic acid moleties, said probe being capable of hybridising to a target sequence of mycobacterial rDNA, precursor rRNA or 23S, 16S or 5S rRNA forming detectable hybrids, and a mixture of such probes.
- 6. Peptide nucleic acid probe according to any one of claims 1 to 5 for detecting a target sequence of rDNA, precursor rRNA or 23S, 16S or 5S rRNA of one or more mycobacteria of the Mycobacterium tuberculosis Complex (MTC) or for detecting a target sequence of rDNA, precursor rRNA or 23S, 16S or 5S rRNA of one or more mycobacteria other than mycobacteria of the Mycobacterium tuberculosis Complex (MOTT) optionally present in a sample, which probe comprises from 10 to 30 polymerised moieties of formula (I)

wherein each X and Y independently designate O or S, each Z independently designates O, S, NR^1 , or $C(R^1)_2$, wherein each R^1 independently designate H, C_{1-6} alkyl, C_{1-6} alkenyl, C_{1-6} alkynyl,

each R², R³ and R⁴ designate independently H, the side chain of a naturally occurring amino acid, the side chain of a non-naturally occurring amino acid, C₁₋₄ alkyl, C₁₋₄ alkenyl or C₁₋₄ alkynyl, or a functional group, each Q independently designates a naturally occurring nucleobase, a non-naturally occurring nucleobase, an intercalator, a nucleobase-binding

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group, a label or H,

with the proviso that the probe comprising such subsequence is capable of forming detectable hybrids with the target sequence of said mycobacterial rDNA, precursor rRNA or 23S, 16S or 5S rRNA.

and a mixture of such probes.

7. Peptide nucleic acid probe according to any one of claims 1 to 6 for detecting a target sequence of 23S rRNA of one or more mycobacteria of the Mycobacterium tuberculosis Complex (MTC) optionally present in a sample, which probe comprises from 10 to 30 polymerised moieties of formula (I) as defined in claim 6.

with the proviso that the Qs of adjacent moieties are selected so as to form a sequence of which a subsequence includes at least one nucleobase that is complementary to a nucleobase of M. tuberculosis 23S rRNA differing from the corresponding nucleobase of at least M. avium located within the following domains

Positions 149-158 in Figure 1A,

Positions 220-221 in Figure 1A.

20 Positions 328-361 in Figure 1A and Figure 1B,

Positions 453-455 in Figure 1B,

Positions 490-501 in Figure 1B,

Positions 637-660 in Figure 1C,

Positions 706-712 in Figure 1D,

25 Positions 762-789 in Figure 1D,

Position 989 in Figure 1D,

Positions 1068-1072 in Figure 1D,

Position 1148 in Figure 1E,

Positions 1311-1329 in Figure 1E.

30 Positions 1361-1364 in Figure 1F,

Position 1418 in Figure 1F,

Positions 1563-1570 in Figure 1F.

Positions 1627-1638 in Figure 1G,

Positions 1675-1677 in Figure 1G,

35 Position 1718 in Figure 1G,

Positions 1734-1740 in Figure 1H,

Positions 1967-1976 in Figure 1H,

Positions 2403-2420 in Figure 1H,

Positions 2457-2488 in Figure 1I,
Positions 2952-2956 in Figure 1I,
Positions 2966-2969 in Figure 1J,
Positions 3000-3003 in Figure 1J or
Positions 3097-3106 in Figure 1J,

and further with the proviso that the probe comprising such subsequence is capable of forming detectable hybrids with a target sequence of said mycobacterial 23S rRNA, and a mixture of such probes.

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8. Peptide nucleic acid probe according to any one of claims 1 to 6 for detecting a target sequence of 16S rRNA of one or more mycobacteria of the Mycobacterium tuberculosis Complex (MTC) optionally present in a sample, which probe comprises from 10 to 30 polymerised moieties of formula (I) as defined in claim 6,

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with the proviso that the Qs of adjacent moieties are selected so as to form a sequence of which a subsequence includes at least one nucleobase that is complementary to a nucleobase of M. tuberculosis 16S rRNA differing from the corresponding nucleobase of at least M. avium located within the following domains

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Positions 76-79 in Figure 2A,
Positions 98-101 in Figure 2A,
Positions 135-136 in Figure 2 A,
Positions 194-201 in Figure 2B,
Positions 222-229 in Figure 2B,
Position 242 in Figure 2B,
Position 474 in Figure 2C,
Positions 1136-1145 in Figure 2C,
Positions 1271-1272 in Figure 2C,
Positions 1287-1292 in Figure 2D,
Position 1313 in Figure 2D, or
Position 1334 in Figure 2D,

and further with the proviso that the probe comprising such subsequence is capable of forming detectable hybrids with a target sequence of said mycobacterial 16S rRNA, and a mixture of such probes.

9. Peptide nucleic acid probe according to any one of claims 1 to 6 for detecting a target

sequence of 5S rRNA of one or more mycobacteria of the Mycobacterium tuberculosis Complex (MTC) optionally present in a sample, which probe comprises from 10 to 30 polymerised moieties of formula (I) as defined in claim 6,

- with the proviso that the Qs of adjacent moieties are selected so as to form a sequence of which a subsequence includes at least one nucleobase that is complementary to a nucleobase of M. tuberculosis 5S rRNA differing from the corresponding nucleobase of at least M. avium located within the following domain
- 10 Positions 86-90 in Figure 3

and further with the proviso that the probe comprising such subsequence is capable of forming detectable hybrids with a target sequence of said mycobacterial 5S rRNA, and a mixture of such probes.

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10. Peptide nucleic acid probe according to any one of claims 1 to 8 for detecting a target sequence of 23S or 16S rRNA of one or more mycobacteria of the Mycobacterium tuberculosis Complex (MTC) optionally present in a sample, which probe comprises from 10 to 30 polymerised moieties of formula (I) as defined in claim 6,

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with the proviso that the Qs of adjacent moieties are selected so as to form a sequence of which a subsequence includes at least one nucleobase that is complementary to a nucleobase of M. tuberculosis 23S or 16 S rRNA differing from the corresponding nucleobase of at least M. avium located within the following domains

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Positions 149-158 in Figure 1A,
Positions 328-361 in Figure 1A and Figure 1B,
Positions 490-501 in Figure 1B,
Positions 637-660 in Figure 1C,
Positions 762-789 in Figure 1D,
Positions 1068-1072 in Figure 1D,
Positions 1311-1329 in Figure 1E,
Positions 1361-1364 in Figure 1F,
Positions 1563-1570 in Figure 1F,
Positions 1627-1638 in Figure 1G,
Positions 1734-1740 in Figure 1H,

Positions 2457-2488 in Figure 1I, Positions 2952-2956 in Figure 1I, Positions 3097-3106 in Figure 1J, Positions 135-136 in Figure 2 A, or Positions 1287-1292 in Figure 2D,

- and further with the proviso that the probe comprising such subsequence is capable of forming detectable hybrids with a target sequence of said mycobacterial 23S or 16S rRNA, and a mixture of such probes.
- 11. Peptide nucleic acid probe according to any one of claims 1 to 6 for detecting a target
 sequence of 23S rRNA of one or more mycobacteria other than mycobacteria of the
 Mycobacterium tuberculosis Complex (MOTT) optionally present in a sample, which probe comprises from 10 to 30 polymerised moieties of formula (I) as defined in claim 6,
- with the proviso that the Qs of adjacent moieties are selected so as to form a sequence of
 which a subsequence includes at least one nucleobase that is complementary to a
 nucleobase of M. avium 23S rRNA differing from the corresponding nucleobase of at least M.
 tuberculosis located within the following domains

Positions 99-101 in Figure 4A,

20 Position 183 in Figure 4A,

Positions 261-271 in Figure 4A,

Positions 281-284 in Figure 4B,

Positions 290-293 in Figure 4B,

Positions 327-335 in Figure 4B,

25 Positions 343-357 in Figure 4B,

Positions 400-405 in Figure 4B and Figure 4C,

Positions 453-462 in Figure 4C,

Positions 587-599 in Figure 4C,

Positions 637-660 in Figure 4D,

30 Positions 704-712 in Figure 4D,

Positions 763-789 in Figure 4E,

Positions 1060-1074 in Figure 4E,

Positions 1177-1185 in Figure 4E,

Positions 1259-1265 in Figure 4F,

35 Positions 1311-1327 in Figure 4F,

Positions 1345-1348 in Figure 4F,

Positions 1361-1364 in Figure 4G,

Positions 1556-1570 in Figure 4G,

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Positions 1608-1613 in Figure 4H,

Positions 1626-1638 in Figure 4H,

Positions 1651-1659 in Figure 4H,

Positions 1675-1677 in Figure 4H.

5 Positions 1734-1741 in Figure 4H,

Positions 1847-1853 in Figure 4I,

Positions 1967-1976 in Figure 4I,

Positions 2006-2010 in Figure 41,

Positions 2025-2027 in Figure 41,

10 Positions 2131-2132 in Figure 4J,

Positions 2252-2255 in Figure 4J,

Positions 2396-2405 in Figure 4J and Figure 4K.

Positions 2416-2420 in Figure 4K,

Positions 2474-2478 in Figure 4K.

15 Position 2687 in Figure 4K,

Position 2719 in Figure 4K,

Position 2809 in Figure 4L,

Positions 3062-2068 in Figure 4L, or

Positions 3097-3106 in Figure 4L,

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and further with the proviso that the probe comprising such subsequence is capable of forming detectable hybrids with a target sequence of said mycobacterial 23S rRNA, and a mixture of such probes.

- 12. Peptide nucleic acid probe according to any one of claims 1 to 6 for detecting a target sequence of 16S rRNA of one or more mycobacteria other than mycobacteria of the Mycobacterium tuberculosis Complex (MOTT) optionally present in a sample, which probe comprises from 10 to 30 polymerised moieties of formula (I) as defined in claim 6,
- with the proviso that the Qs of adjacent moieties are selected so as to form a sequence of which a subsequence includes at least one nucleobase that is complementary to a nucleobase of M. avium 16S rRNA differing from the corresponding nucleobase of at least M. tuberculosis located within the following domains
- Positions 135-136 in Figure 5A,
 Positions 472-475 in Figure 5A,
 Positions 1136-1144 in Figure 5A,
 Positions 1287-1292 in Figure 5B,

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Position 1313 in Figure 5B, or Position 1334 in Figure 5B,

and further with the proviso that the probe comprising such subsequence is capable of forming
 detectable hybrids with a target sequence of said mycobacterial 16S rRNA,
 and a mixture of such probes.

13. Peptide nucleic acid probe according to any one of claims 1 to 6, 11 and 12 for detecting a target sequence of 23S or 16S rRNA of one or more mycobacteria other than mycobacteria of the Mycobacterium tuberculosis Complex (MOTT) optionally present in a sample, which probe comprises from 10 to 30 polymerised moleties of formula (I) as defined in claim 6,

with the proviso that the Qs of adjacent moieties are selected so as to form a sequence of which a subsequence includes at least one nucleobase that is complementary to a nucleobase of M. avium 23S or 16S rRNA differing from the corresponding nucleobase of at least M. tuberculosis located within the following domains

Positions 99-101 in Figure 4A, Positions 290-293 in Figure 4B,

20 Positions 400-405 in Figure 4B and Figure 4C.

Positions 453-462 in Figure 4C,

Positions 637-660 in Figure 4D,

Positions 763-789 in Figure 4E,

Positions 1311-1327 in Figure 4F,

25 Positions 1361-1364 in Figure 4G,

Positions 1734-1741 in Figure 4H,

Positions 2025-2027 in Figure 4I,

Positions 2474-2478 in Figure 4K,

Positions 3062-2068 in Figure 4L, or

30 Positions 1287-1292 in Figure 5B,

and further with the proviso that the probe comprising such subsequence is capable of forming detectable hybrids with a target sequence of said mycobacterial 23S or 16S rRNA, and a mixture of such probes.

14. Peptide nucleic acid probe according to any one of claims 1 to 6 for detecting a target sequence of 23S, 16S or 5S rRNA of one or more mycobacteria of the Mycobacterium tuberculosis Complex (MTC) or for detecting a target sequence of 23S, 16S or 5S rRNA of

one or more mycobacteria other than mycobacteria of the Mycobacterium tuberculosis Complex (MOTT) optionally present in a sample, which probe comprises from 10 to 30 polymerised moleties of formula (I) as defined in claim 6,

- with the proviso that the Qs of adjacent moieties are selected so as to form a sequence of which a subsequence includes at least one nucleobase that is complementary to a nucleobase that differs from the corresponding nucleobase of 23S, 16S or 5S rRNA of said one or more mycobacteria located within the following domains
- 10 positions 2568-2569 in Figure 6,

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Position 452 in Figure 7, Positions 473-477 in Figure 7, or Positions 865-866 in Figure 7,

and further with the proviso that the probe comprising such subsequence is capable of forming detectable hybrids with the target sequence of said mycobacterial 23S, 16S or 5S rRNA, and a mixture of such probes.

15. Peptide nucleic acid probe according to any one of claims 1 to 14 of formula (II), (III), or (IV)

$$\mathbb{Z}^{\mathbb{Z}^2}$$

25 (II)

wherein Z, R^2 , R^3 , and R^4 , and Q is as defined in claim 6 with the provisos defined in claims 6

to 14,

and a mixture of such probes.

16. Peptide nucleic acid probe according to any one of claims 1 to 15, wherein Z is NH, NCH₃ or O, each R², R³ and R⁴ independently designate H or the side chain of a naturally occurring amino acid, the side chain of a non-naturally occurring amino acid, or C₁₋₄ alkyl, and each Q is a naturally occurring nucleobase or a non-naturally occurring nucleobase with the provisos defined in claims 6 to 14,

and a mixture of such probes.

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- 17. Peptide nucleic acid probe according to any one of claims 1 to 16, wherein Z is NH or O, and R² is H or the side chain of Ala, Asp, Cys, Glu, His, HomoCys, Lys, Orn, Ser or Thr, and Q is a nucleobase selected from thymine, adenine, cytosine, guanine, uracil, iso-C and 2,6-diaminopurine with the provisos defined in claims 6 to 14,
- 15 and a mixture of such probes.
 - 18. Peptide nucleic acid probe according to any one of claims 1 to 17 of formula (V)

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wherein R⁴ is H or the side chain of Ala, Asp, Cys, Glu, His, HomoCys, Lys, Orn, Ser or Thr, and Q is as defined in claim 17 with the provisos defined in claims 6 to 14, and a mixture of such probes.

- 25
- 19. Peptide nucleic acid probe according to any one of claims 1 to 18 further comprising one or more labels and a mixture of such probes, which labels may be mutually identical or different, which probes optionally may comprise one or more linkers, and which probes may be mutually identical or different with the provisos defined in claims 6 to 14.
- 30
- 20. Peptide nucleic acid probe according to any one of claims 1 to 19 for detecting a target sequence of one or more mycobacteria, the nucleobase sequence of said probe being substantially complementary to the nucleobase sequence of said target sequence.

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21. Peptide nucleic acid probe according to any one of claims 1 to 20 for detecting a target sequence of one or more mycobacteria, the nucleobase sequence of said probe being complementary to the nucleobase sequence of said target sequence.

22. Peptide nucleic acid probes according to any one of claims 1 to 21, wherein the Qs of adjacent moieties are selected so as to form the following subsequences

5	AGA TGC GGG TAG CAC (selected from positions 149-158 in Figure 1A),	(Seq ID no 1)
	TGT TTT CTC CTC CTA (selected from positions 220-221 in Figure 1A),	(Seq ID no 2)
	ACT GCC TCT CAG CCG (selected from positions 328-361 in	
	Figure 1A and Figure 1B),	(Seq ID no 3)
	TGA TAC TAG GCA GGT (selected from positions 453-455 in Figure 1B),	(Seq ID no 4)
10	CGG ATT CAC AGC GGA (selected from positions 490-501 in Figure 1B),	(Seq ID no 5)
	TCA CCA CCC TCC TCC (selected from positions 637-660 in Figure 1C),	(Seq ID no 6)
	TTA ACC TTG CGA CAT (selected from positions 706-712 in Figure 1C),	(Seq ID no 7)
	ACT ATT CAC ACG CGC (selected from positions 762-789 in Figure 1D),	(Seq ID no 8)
	CTC CGC GGT GAA CCA (selected from position 989 in Figure 1D),	(Seq ID no 9)
15	GCT TTA CAC CAC GGC (selected from positions 1068-1072 in Figure 1D),	(Seq ID no 10)
	ACG CTT GGG GGC CTT (selected from position 1148 in Figure 1E),	(Seq ID no 11)
	CCA CAC CCA CCA CAA (selected from positions 1311-1329 in Figure 1E),	(Seq ID no 12)
	CCG GTG GCT TCG CTG (selected from positions 1361-1364 in Figure 1F),	(Seq ID no 13)
	ACT TGC CTT GTC GCT (selected from position 1418 in Figure 1F),	(Seq ID no 14)
20	GAT TCG TCA CGG GCG (selected from positions 1563-1570 in Figure 1F),	(Seq ID no 15)
	AAC TCC ACA CCC CCG (selected from positions 1627-1638 in Figure 1G),	(Seq ID no 16)
	ACT CCA CAC CCC CGA (selected from positions 1627-1638 in Figure 1G),	(Seq ID no 17)
	ACC CCT TCG CTT GAC (selected from positions 1675-1677 in Figure 1G),	(Seq ID no 18)
	CTT GCC CCA GTG TTA (selected from position 1718 in Figure 1G),	(Seq ID no 19)
25	CTC TCC CTA CCG GCT (selected from positions 1734-1740 in Figure 1H),	(Seq ID no 20)
	GAT ATT CCG GTC CCC (selected from positions 1967-1976 in Figure 1H),	(Seq ID no 21)
	ACT CCG CCC CAA CTG (selected from positions 2403-2420 in Figure 1H),	(Seq ID no 22)
	CTG TCC CTA AAC CCG (selected from positions 2457-2488 in Figure 1I),	(Seq ID no 23)
	TTC GAG GTT AGA TGC (selected from positions 2457-2488 in Figure 1I),	(Seq ID no 24)
30	GTC CCT AAA CCC GAT (selected from positions 2457-2488 in Figure 1I),	(Seq ID no 25)
	GGT GCA CCA GAG GTT (selected from positions 2952-2956 in Figure 1I),	(Seq ID no 26)
	CTG GCG GGA CAA CTG (selected from positions 2966-2969 in Figure 1J),	(Seq ID no 27)
	TTA TCC TGA CCG AAC (selected from positions 3000-3003 in Figure 1J),	(Seq ID no 28)
	GAC CTA TTG AAC CCG (selected from positions 3097-3106 in Figure 1J),	(Seq ID no 29)
35		
	GAA GAG ACC TTT CCG (selected from positions 76-79 in Figure 2A),	(Seq ID no 30)
	CAC TCG AGT ATC TCC (selected from positions 98-101 in Figure 2A),	(Seq ID no 31)
	ATC ACC CAC GTG TTA (selected from positions 136-136 in Figure 2A),	(Seq ID no 32)
	GCA TCC CGT GGT CCT (selected from positions 194-201 in Figure 2B),	(Seq ID no 33)
40	CAC AAG ACA TGC ATC (selected from positions 194-201 in Figure 2B),	(Seq ID no 34)
	TAA AGC GCT TTC CAC (selected from positions 222-229 in Figure 2B),	(Seq ID no 35)
	GCT CAT CCC ACA CCG (selected from position 242 in Figure 2B),	(Seq ID no 36)

GAG ACA GTT GGG AAG (selected from positions 2252-2255 in Figure 4J),

TGG CGT CTG TGC TTC (selected from positions 2396-2405 in

(Seq ID no 74)

	Figure 4J and Figure 4K),	(Seq ID no 75)
	CGA CTC CAC ACA AAC (selected from positions 2416-2420 in Figure 4K),	(Seq ID no 76)
	GAT AAG GGT TCG ACG (selected from positions 2474-2478 in Figure 4K),	(Seq ID no 77)
	ATC CGT TGA GTG ACA (selected from position 2687 in Figure 4K),	(Seq ID no 78)
5	CAG CCC GTT ATC CCC (selected from position 2719 in Figure 4K),	(Seq ID no 79)
	AAC CTT TGG GAC CTG (selected from position 2809 in Figure 4L),	(Seq ID no 80)
	TAA AAG GGT GAG AAA (selected from positions 3062-3068 in Figure 4L),	(Seq ID no 81)
	GTC TGG CCT ATC AAT (selected from positions 3097-3106 in Figure 4L),	(Seq ID no 82)
	·	•
10	AGA TTG CCC ACG TGT (selected from positions 135-136 in Figure 5A),	(Seq ID no 83)
	AAT CCG AGA AAA CCC (selected from positions 472-475 in Figure 5A),	(Seq ID no 84)
	GCA TTA CCC GCT GGC (selected from positions 1136-1144 in Figure 5B),	(Seq ID no 85)
	TTA AAA GGA TTC GCT (selected from positions 1287-1292 in Figure 5B),	(Seq ID no 86)
	AGA CCC CAA TCC GAA (selected from position 1313 in Figure 5B),	(Seq ID no 87)
15	GAC TCC GAC TTC ATG (selected from position 1334 in Figure 5B),	(Seq ID no 88)
	GTC TTT TCG TCC TGC (selected from positions 2568-2569 in Figure 6),	(Seq ID no 89)
	GTC TTA TCG TCC TGC (selected from positions 2568 in Figure 6),	(Seq ID no 90)
	GTC TTC TCC TCC (selected from positions 2568 in Figure 6),	(Seq ID no 91)
20	GTC TTG TCG TCC TGC (selected from positions 2568 in Figure 6),	(Seq ID no 92)
	GTC TAT TCG TCC TGC (selected from positions 2568 in Figure 6),	(Seq ID no 93)
	GTC TCT TCG TCC TGC (selected from positions 2568 in Figure 6),	(Seq ID no 94)
	GTC TGT TCG TCC TGC (selected from positions 2568 in Figure 6),	(Seq ID no 95)
25	TTG GCC GGT GCT TCT (selected from positions 452 in Figure 7),	(Seq ID no 96)
	TTG GCC GGT ACT TCT (selected from positions 452 in Figure 7).	(Seq ID no 97)
	TTG GCC GGT CCT TCT (selected from positions 452 in Figure 7),	(Seq ID no 98)
•	TTG GCC GGT TCT TCT (selected from positions 452 in Figure 7),	(Seq ID no 99)
	ACC GCG GCT GCT GGC (selected from positions 473-477 in Figure 7),	(Seq ID no 100)
30	ACC GCG GCT ACT GGC (selected from positions 473 in Figure 7),	(Seq ID no 101)
	ACC GCG GCT CCT GGC (selected from positions 473 in Figure 7), or	(Seq ID no 102)
	ACC GCG GCT TCT GGC (selected from positions 473 in Figure 7),	(Seq ID no 103)
	CGG CAG CTG GCA CGT (selected from positions 474 in Figure 7),	(Seq ID no 104)
	CGG CCG CTG GCA CGT (selected from positions 474 in Figure 7),	(Seq ID no 105)
35	CGG CTG CTG GCA CGT (selected from positions 474 in Figure 7),	(Seq ID no 106)
	CGT ATT ACC GCA GCT (selected from positions 477 in Figure 7),	(Seq ID no 107)
	CGT ATT ACC GCC GCT (selected from positions 477 in Figure 7),	(Seq ID no 107)
	CGT ATT ACC GCT GCT (selected from positions 477 in Figure 7),	(Seq ID no 109)
	TTC CTT TGA GTT TTA (selected from positions 865-866 in Figure 7),	(Seq ID no 110)
40	TTC CTT TAA GTT TTA (selected from positions 865 in Figure 7),	(Seq ID no 111)
	TTC CTT TCA GTT TTA (selected from positions 865 in Figure 7),	(Seq ID no 112)
	TTC CTT TTA GTT TTA (selected from positions 865 in Figure 7),	(Seq ID no 113)
	TTC CTT AGA GTT TTA (selected from positions 866 in Figure 7),	(Seq ID no 114)

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•	68	
TTC CTT CGA GTT TTA (se	elected from positions 866 in Figure 7),	(Seq ID no 115)
TTC CTT GGA GTT TTA (se	elected from positions 866 in Figure 7),	(Seq ID no 116)
CAT GTG TCC TGT GGT		(Seq ID no 117)
CGT CAG CCC GAG AAA		(Seg ID no 118)
CAC TAC ACA CGC TCG		(Seg ID no 119)
TGG CGT TGA GGT TTC a	nd	(Seg ID no 120)
AAC ACT CCC TTT GGA		(Seq ID no 123)
and a mixture of such pro	oes.	

23. Peptide nucleic acid probes according to claim 22, wherein the Qs of adjacent moieties are selected so as to form the following subsequences

	TCA CCA CCC TCC TCC	(Seq ID no 6)
15	CCA CCC TCC TCC	(modified Seq ID no 6)
	ACT ATT CAC ACG CGC	(Seq ID no 8)
	CCA CAC CCA CCA CAA	(Seq ID no 12)
	AAC TCC ACA CCC CCG	(Seq ID no 16)
	ACT CCA CAC CCC CGA	(Seq ID no 17)
20	ACT CCG CCC CAA CTG	(Seq ID no 22)
	CTG TCC CTA AAC CCG	(Seq ID no 23)
	TTC GAG GTT AGA TGC	(Seq ID no 24)
	GTC CCT AAA CCC GAT	(Seq ID no 25)
	GAC CTA TTG AAC CCG	(Seq ID no 29)
25		
	GCA TCC CGT GGT CCT	(Seq ID no 33)
	CAC AAG ACA TGC ATC	(Seg ID no 34)
	GGC TTT TAA GGA TTC	(Seq ID no 40)
30	GAT CAA TGC TCG GTT	(Seq ID no 44)
	CGA CTC CAC ACA AAC	(Seq ID no 76)
	GCA TTA CCC GCT GGC	(Seq ID no 85)
		·
35	GTC TTA TCG TCC TGC	(Seq ID no 90)
	GTC TTC TCG TCC TGC	(Seq ID no 91)
	GTC TTG TCG TCC TGC	(Seq ID no 92)
	GTC TAT TCG TCC TGC	(Seq ID no 93)
	GTC TCT TCG TCC TGC	(Seq ID no 94)
40	GTC TGT TCG TCC TGC	(Seq ID no 95)
	AAC ACT CCC TTT GGA	(Seq ID no 123)

CAT GTG TCC TGT GGT	(Seq ID no 117)
CGT CAG CCC GAG AAA	(Seq ID no 118)
CAC TAC ACA CGC TCG, TGG CGT TGA GGT TTC	(Seq ID no 119) (Seq ID no 120)

and a mixture of such probes.

10 24. Peptide nucleic acid probes according to claim 22 or 23 selected from

	Lys(Flu)-Lys(Flu)-TCA CCA CCC TCC TCC-NH2	(OK 446/modified Seq ID no 6)
	Lys(Flu)-Lys(Flu)-CCA CCC TCC TCC-NH₂	(OK 575/modified Seq ID no 6)
	Lys(Flu)-Lys(Flu)-ACT ATT CAC ACG CGC-NH₂	(OK 447/modified Seq ID no 8)
15	Lys(Flu)-ACT ATT CAC ACG CGC-NH ₂	(OK 688/modified Seq ID no 8)
	Lys(Flu)-Lys(Flu)-CCA CAC CCA CCA CAA-NH2	(OK 448/modified Seq ID no 12)
	Lys(Flu)-Lys(Flu)-AAC TCC ACA CCC CCG-NH2	(OK 449/modified Seq ID no 16)
	Lys(Flu)-Lys(Flu)-ACT CCA CAC CCC CGA-NH ₂	(OK 309/modified Seq ID no 17)
	Lys(Flu)-Lys(Flu)-ACT CCG CCC CAA CTG-NH2	(OK 450/modified Seq ID no 22)
20	Lys(Flu)-Lys(Flu)-CTG TCC CTA AAC CCG-NH₂	(OK 305/modified Seq ID no 23)
	Lys(Flu)-Lys(Flu)-TTC GAG GTT AGA TGC-NH₂	(OK 306/modified Seq ID no 24)
	Lys(Flu)-TTC GAG GTT AGA TGC-NH₂	(OK 682/modified Seq ID no 24)
	Lys(Flu)-Lys(Flu)-GTC CCT AAA CCC GAT-NH₂	(OK 307/modified Seq ID no 25)
	Lys(Flu)-GTC CCT AAA CCC GAT-NH₂	(OK 654/modified Seq ID no 25)
25	Lys(Flu)-GAC CTA TTG AAC CCG-NH₂	(OK 660/modified Seq ID no 29)
		·
	Lys(Flu)-Lys(Flu)-Gly-GCA TCC CGT GGT CCT-NH₂	(OK 223/modified Seq ID no 33)
	Lys(Flu)-Lys(Flu)-CAC AAG ACA TGC ATC-NH₂	(OK 310/modified Seq ID no 34)
	Lys(Flu)-CAC AAG ACA TGC ATC-NH₂	(OK 655/modified Seq ID no 34)
30	Lys(Flu)-GGC TTT TAA GGA TTC-NH₂	(OK 689/modified Seq ID no 40)
	Lys(Rho)-GGC TTT TAA GGA TTC-NH₂	(OK 702/modified Seq ID no 40)
	Flu-β-Ala-β-Ala-GAT CAA TGC TCG GTT-NH ₂	(OK 624/modified Seq ID no 44)
	Flu-β-Ala-β-Ala-CGA CTC CAC ACA AAC-NH ₂	(OK 612/modified Seq ID no 76)
35		
	Flu-β-Ala-β-Ala-GCA TTA CCC GCT GGC-NH ₂	(OK 623/modified Seq ID no 85)
	•	(**************************************
	Lys(Flu)-GTC TTT TCG TCC TGC-NH₂	(OK 745/modified Seq ID no 89)
	Lys(Rho)-GTC TTA TCG TCC TGC-NH ₂	(OK 746/modified Seq ID no 90)
40	Lys(Rho)-GTC TTC TCG TCC TGC-NH ₂	(OK 746/modified Seq ID no 91)
	Lys(Rho)-GTC TTG TCG TCC TGC-NH ₂	(OK 746/modified Seq ID no 92)
	Lys(Rho)-GTC TAT TCG TCC TGC-NH ₂	(OK 747/modified Seq ID no 93)
	-	,

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Lys(Rho)-GTC TCT TCG TCC TGC-NH₂ Lys(Rho)-GTC TGT TCG TCC TGC-NH₂ (OK 747/modified Seq ID no 94) (OK 747/modified Seq ID no 95)

Lys(Flu)-AAC ACT CCC TTT GGA-NH2

(OK 749/modified Seq ID no 123)

5

wherein Flu denotes a 5-(and 6)-carboxyfluoroescein label and Rho denotes a modamine label,

and a mixture of such probes.

- 10 25. Use of a peptide nucleic acid probe according to any one of claims 1 to 24 or a mixture thereof for detecting a target sequence of one or more mycobacteria optionally present in a sample.
- 26. Use of a peptide nucleic acid probe or a mixture thereof according to claim 25 for detecting
 a target sequence of one or more mycobacteria of the Mycobacterium tuberculosis Complex
 (MTC), in particular a target sequence of M. tuberculosis.
 - 27. Use of a peptide nucleic acid probe or a mixture thereof according to claims 25 for detecting a target sequence of one or more mycobacteria other than mycobacteria of the Mycobacterium tuberculosis Complex, in particular a target sequence of one or more mycobacteria of the Mycobacterium avium Complex.
 - 28. Method for detecting a target sequence of one or more mycobacteria optionally present in a sample comprising

25

(1) contacting any rRNA or rDNA present in said sample with one or more peptide nucleic acid probes according to any one of claims 1 to 24 or a mixture thereof under conditions, whereby hybridisation takes place between said probe(s) and said rRNA or rDNA, and

30

- (2) observing or measuring any formed detectable hybrids, and relating said observation or measurement to the presence of a target sequence of one or more mycobacteria in said sample.
- 35 29. Method according to claim 28 for detecting a target sequence of one or more mycobacteria of the Mycobacterium tuberculosis Complex (MTC), in particular a target sequence of M. tuberculosis.
 - 30. Method according to claim 28 for detecting a target sequence of one or more

5

mycobacteria other than mycobacteria of the Mycobacterium tuberculosis Complex.

- 31. Method according to any one of claims 28 to 30, wherein the hybridisation takes place in situ.
- 32. Method according to any of of claims 28 to 30, wherein the hybridisation takes place in vitro.
- 33. A method according to any one of claims 28 to 32,
- 10 characterised in that a signal amplifying system is used for measuring the resulting hybridisation.
 - 34. Method according to any one of claims 28 to 33, wherein the sample is a sputum sample.
- 35. Kit for detecting a target sequence of one or more mycobacteria, in particular a target sequence of one or more mycobacteria of the Mycobacterium tuberculosis Complex (MTC), in particular a target sequence of M. tuberculosis, and/or for detecting a target sequence of one or more mycobacteria other than mycobacteria of the Mycobacterium tuberculosis Complex (MOTT), in particular a target sequence of one or more mycobacteria of the Mycobacterium avium Complex.
 - c h a r a c t e r i s e d in that said kit comprises at least one peptide nucleic acid probe according to any one of claims 1 to 24, and optionally a detection system with at least one detecting reagent.
- 36. Kit according to claim 35,c h a r a c t e r i s e d in that it further comprises a solid phase capture system.

		130	140	150	16	0
1093	GGGGAAAG	CCAGCACGA	GTGATGTCGT	CTACCCGCA	re r	M.tuberculosis
422	GGGGAAG	CCCAGCACGA	STGATGTCGT	TTACCCGTA	TCT.	M. avium
422	GGGGGAAA	CCAGCACGAC	こかにかからかってきから		n C m	M. managerile
507	GGGGGAAG	CCEGCACGA	STGATGTCGT	DAGCCDACT	3Cm	M phlai
432	GGGGAAA	CCAACACGA	STEATGTCGTO STEADGTCGTO STEATGTCGTO	THE CCCCIDE		M lenne
207	GGGGAAAC	CCAGCACGA	STRATGTCGTG	SHIPACCCGHA	וייים.	M. doctni
150	GGGGAAAG	CCAGCACGA	3TGBTGTCGTG		ייריי	M.kansasii
2588	GGGGAAAC	CCGGCACGAC	37671676676 37677676676	TOPCOOCK	300	M.smegmatis
		3 COE O 11 CO 11 C	JIONIGIOGIC	ST OUCCHOPCE	301	M. Smegmatis
						•
				•		
		210	220	230	240	
1172	CATCTCAG	TACCCGTAG	FIGGAGAAAAC	AATTGTGATT	rcc	M.tuberculosis
501	CATCTCAC	TACCCGTAGE	CACACANANC	ים תבות בית בית תולי	חככ	M orrive
501	CATCTCAG	TACCCGTAG	AGAAGAAAAC	AATTGTGATT	rcc	M.paratuberc. M.phlei M.leprae
586	CATCTCAG	TACCCGTAG	AGAGAAAAC	ביים ביים ביים ביים	יככ	M phlei
511	CATCTCAG	TACCCGTAGE	AGAGAAAAC	בנב בכונה. השמשה השמש בני	ירכ	M lenne
286	CATCTCAG	TACCCGTAGE	AGAAGAAAAC	ייייעטינטעסעמיי	יככ	M. reptae
229	CATCTCAG	TACCCGTAGE	BEBBEBBBB	'A DE A CTCATT	יככ	M.kansasii
	CATCTCAG	TECCGTAGE	PAGAGAAAAC	ANDROTOATI	יככ	M.smegmatis
	0.1101010	.1[]00001M00	AND	WHITGIGHT		M.Smegmatis
				1		
		· 				
		330	340	350	360	•
1000						
1289	TGTGGGAG	GAVATETO	CAGCGCTACC	CGGCTGAGA-	GG	M.tuberculosis
617	TGTGGGAT	'TGATATGTC'I	CAGOTCTACC	IGGCTGAGG-	GG	M.avium
617	TGTGGGAT	<u>TGATATGTCI</u>	CAGCIICTACC	<u>TGGCTGAGG</u> -	GG	M.paratuberc.
703	TGTGGGGC	CTGTGTGTC-	CATCGTCCGC	CGGCGATGGC	AG	M.phlei
629	TGTGGGAT	TGGTATGTCI	CAACIICTACC	NGGNTGAGG-	GG	M lenrae
404	TGTGGGAIT	CGATACGTCT	CAGCITCTACC	CGGCTGAGG	CC	M coetri
347	TGTGGGAT	'QGATAQGTC'I	CAGCIICTACC	CGGCTGAGG	·GG	M.kansasii
2785	TGTGGGAC	CTATOTUTC	CGCCTCTACC	TGGCTGFGAG	GG	M.smegmatis
			-		•	3

Figure 1A

	;	370	380	390	400
1327	CAGTCAGA	AAGTGTCGT	GTTAGCGGA	AGTECCTECE	AT M.tuberculosis
656	HAGTCAGA	AAGTGTCGT	GTTAGCGGA	AGTGGCCTGGG	AD M.avium
656	HAGTCAGA	AAGTGTCGT	GTTDGCGGD	AGTGGCCTGGG	AC M.paratuberc.
742	TAGTGATIA	AAGCAGTGT	GTTAGGTGA	AGTGGCCTGGG	AG M. paracuberc.
668	TAGTCAGAZ	ABGTGCGTG	301130 <u>01</u> 030	Prescriss	AT M.leprae
443	CAGTCAGA	¥₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽	SGTTADCGGA:	JG1166CC1666	AT M.Teprae AT M.gastri
386	CAGTCAGA	1701010010 180707070070	CTTANCCCA	GTGGCCTGGG.	
2823	CAGTGAGA	2201010010 2200000000000000000000000000	COUNT CCCVI	A TGGCCTGGG	AT M.kansasii
2025	OVOTPLOY	- ANDIGITATION	3G I TAGCGGA	Mareechiege	AT M.smegmatis
				•	·
		- 			
		150	460	_	480
1406	CGGCACCT	CCTAGTATO	CAATTCCCGAG	TAGCAGCGGG	CC M.tuberculosis
735	CGGCACCTG	CCTTATATO	CAACACCCGAG	TAGCAGCGGG	CC M.avium
735	CGGCACCTG	CCTTATATO	CAACACCCGAG	TAGCAGCGGG	CC M.paratuberc
820	TGCTGCC-C	FCTGTCACAG	GTCCCGAG	TAGCAGCGGG	CC M.phlei
747	IGGCACCTG	CCTTGTATO	CAATTCCCGAG	TAGCAGCGGG	CC M.leprae
522	CGGCACCTG	CCTTGTATO	AATTCCCGAG	TAGCAGCGGG	CC M.gastri
465	CGGCACCTG	CCTTGTATC	AATTCCCGAG	TAGCAGCGGG	CC M.kansasii
2902	CGACGTCTG	TCTTGATGG	TGTTCCCGAG	TAGCAGCGGG	CC M.smegmatis
					
	4	190	500	510	520
1446	CGTGGAATC	CGCTGTGA	TCCGCGGA	CCACCGGTA	AG M.tuberculosis
775	CGTGGAATC	AIGCTGTGAP	TCIGCCGGGA	CCACCCGGTAI	AG M.avium
775	CGTGGAATC	TGCTGTGAP	TCTGCCGGGA	CCACCCGGTA	AG M.paratuberc.
857	CGTGGAATC	IGCTGTGAA	TCTGCCGGGA	CCACCCGGTA	AG M.phlei
787	CGTGGAATC	TGCTGTGAA	TCTGCCGGGA	CCACCCGGTA	AG M.leprae
562	CGTGGAATC	IGCTGTGAA	TCTGCCGGGA	CCACCCGGTAZ	AG M.gastri
505	CGTGGAATC	TGCTGTGAA	TOTGCCGGGA	CCACCCGGTA	AG M.kansasii
2942	CGTGGAATC	TGCTGTGAA	TCIGCCGGGA	CCACCCGGTA	AG M.smegmatis
-			- 20000		- m. smegmatts

Figure 1B

		610	620	630	640
566	GTACCTG	AAACCGT	GTGCCTACAAT	CCGTCAGAG	COTCCT M.tuberculos
94	GTACCTG	BAAACCGT	GTGCCTACAAT	CCGTCAGAG	CCTCCT M.avium
94	GTACCTG	BAAACCGT	GTGCCTACAAT	CCGTCAGAG	CCTCCT M.paratuberc
76	GTACCTG	AAACCGT	GTGCCTACAAT	CCGTCADAG	CCCTCT M.phlei
07	GTACCTG	HAAACCGT	GTGCCTACAA1	CCGTCAGAG	CCTCTT M lenne
82	GTACCTG	AAACCGT	GTGCCTACAAT	CCGTCAGAG	CCCTT M.gastri
25	GTACCTG	BAAACCGT	GTGCCTACAAT	CCGTCAGAG	COCTUTE M Kangagii
062	GTACCTG	AAACCGT	GGGC[[TACAAT	СССТСДСДС	CCCTCG M.smegmatis
				OOUTOAGAG.	COLICE M. Smegmatis
		650	660	670	680
606	TTTCCTC	TCCGGAG	SAGGGTEGTGA	TGGCGTGCC	TTTTGA M.tuberculosi
34	C		GIGGGGTGA	TGGCGTGCC	TTTTGA M.avium
34	C		GIGGGGTGA	TGGCGTGCC	TTTTGA M.paratuberc.
016	CTT	G	iagigeetea	TGGCGTGCC	TTTTGA M.phlei
47	T		- GIGGGGTGA	TGGCGTGCI	TTTTGA M.leprae
22	T		GIGGGGTGA	reecerecci	TTTTGA M.gastri
65	c		GIGGGGTGA	TGGCGTGCCT	TTTTGA M.kansasii
102	ACGTGT-				
	PC0101		GIIGGGGTGA	TGGCGTGCC1	TTTTGA M.smegmatis
	ACO101	·	GIIGGGGTGA	TGGCGTGCC1	FTTTGA M.smegmatis
		690	- pi		····
		690	700	710	720
	AGAATGA	GCCTGCGA	700 AGTCAGGGACA	710	720
546	AGAATGA:	GCCTGCGF GCCTGCGF	700 AGTCAGGGACA	710 TGTCGCAAGG	720 GTTAAC M.tuberculosi
546 59	AGAATGA AGAATGA AGAATGA	GCCTGCGA GCCTGCGA GCCTGCGA	700 AGTCAGGGACA AGTCAGGGACA	710 TGTCGCAAGG TGTCGCAAGG	720 GTTAAC M.tuberculosi GTTAAC M.bovis
546 59 3	AGAATGA AGAATGA AGAATGA AGAATGA	GCCTGCGF GCCTGCGF GCCTGCGF	700 AGTCAGGGACA AGTCAGGGACA AGTCAGGGACA	710 TGTCGCAAGG TGTCGCAAGG GTCGCGAGG	720 GTTAAC M.tuberculosi GTTAAC M.bovis GTTAAC M.avium
646 59 3	AGAATGA AGAATGA AGAATGA AGAATGA	GCCTGCGF GCCTGCGF GCCTGCGF GCCTGCGF	700 AGTCAGGGACA AGTCAGGGACA AGTCAGGGACA AGTCAGGGACA	710 TGTCGCAAGG TGTCGCAAGG GTCGCGAGG	720 GTTAAC M.tuberculosi GTTAAC M.bovis GTTAAC M.avium GTTAAC M.intracellul
646 59 3 59	AGAATGA AGAATGA AGAATGA AGAATGA AGAATGA	GCCTGCGA GCCTGCGA GCCTGCGA GCCTGCGA GCCTGCGA	700 AGTCAGGGACA AGTCAGGGACA AGTCAGGGACA AGTCAGGGACA	710 TETCECAAGG TETCECAAGG ETCECEAGG ETCECEAGG	720 GTTAAC M.tuberculosi GTTAAC M.bovis GTTAAC M.avium GTTAAC M.intracellul GTTAAC M.paratuberc.
646 59 3 59 046 72	AGAATGA AGAATGA AGAATGA AGAATGA AGAATGA AGAATGA	GCCTGCGF GCCTGCGF GCCTGCGF GCCTGCGF GCCTGCGF GCCTGCGF	700 AGTCAGGGACA AGTCAGGGACA AGTCAGGGACA AGTCAGGGACA AGTCAGGGACA AGTCAGGGACA	710 TETCECA AGG TETCECA AGG ETCECEA GG ETCECEA GG ETCECEA GG ETCECEA GG TETCECEA GG	720 GTTAAC M.tuberculosi GTTAAC M.bovis GTTAAC M.avium GTTAAC M.intracellul GTTAAC M.paratuberc. GTTAAC M.phlei
646 59 3 59 046 72 47	AGAATGA AGAATGA AGAATGA AGAATGA AGAATGA AGAATGA AGAATGA	GCCTGCGF GCCTGCGF GCCTGCGF GCCTGCGF GCCTGCGF GCCTGCGF GCCTGCGF	700 AGTCAGGGACA AGTCAGGGACA AGTCAGGGACA AGTCAGGGACA AGTCAGGGACA AGTCAGGGACA AGTCAGGGACA	710 TETCECA AGE	720 GTTAAC M.tuberculosi GTTAAC M.bovis GTTAAC M.avium GTTAAC M.intracellul GTTAAC M.paratuberc. GTTAAC M.phlei GTTAAC M.leprae GTTAAC M.gastri
646 59 3 59 046 72 47	AGAATGA AGAATGA AGAATGA AGAATGA AGAATGA AGAATGA AGAATGA AGAATGA	GCCTGCGP GCCTGCGP GCCTGCGP GCCTGCGP GCCTGCGP GCCTGCGP GCCTGCGP GCCTGCGP	700 AGTCAGGGACA AGTCAGGGACA AGTCAGGGACA AGTCAGGGACA AGTCAGGGACA AGTCAGGGACA AGTCAGGGACA AGTCAGGGACA	710 TETCECA AGG TETCECA AGG ETCECEA AGG ETCECEA AGG TETCECEA AGG TETCECEA AGG TETCECEA AGG TETCECEA AGG	720 GTTAAC M.tuberculosi GTTAAC M.bovis GTTAAC M.avium GTTAAC M.intracellul GTTAAC M.paratuberc.

Figure 1C

	770	780	790	800
84 CGACCCI 1039 CG 1039 CG 1126 CGTATCI 1052 CGTAT 827 CGTAT 770 CGTAT	ACACGCGCA1 -CATCCCTTT -CATCCCTTTT -CATCCTTTTG -CACGTGTGA -CACGCGTGA -CACGCGTGA -CACGCGTGA -CGCGCGCGA	FACGCGCGTGT FACGCGCGTGT FTGGGGTGT FTGGGGTTGGTGT FGGGGTTGGTGT FGCGTGTGT FGCGTGTGT	FGAATAGTGG FGAATAGTGG FGAGTGG FGAGTGG FGAGTGG FGAGTGG FGAGTGG	CGTGT M.tuberculo CGTGT M.bovis CGTGT M.avium CGTGT M.intracell CGTGT M.paratuber TGTGT M.phlei CGTGT M.leprae CGTGT M.gastri CGTGT M.kansasii

	970	980	990	1000	
1926	ATTTAGGTGCAGCG'	TGCGTGGTTC	ACCGCGGAGG	TAGAG M.tuk	erculosis
1228	ATTTAGGTGCAGCG	TGCGTGGTTC	ACCACGGAGG	TAGAG M.avi	11m
1228	ATTTAGGTGCAGCG	TGCGTGGTTC	ACCACGGAGG	TAGAG M.par	atuberc.
1322	ATTTAGGTGCAGCGT	rdgcarghttc	TTATCGGAGG	TAGAG M. nhl	ei
1244	ATTTAGGTGCAGCGT	TGCGTGGTTC	ACCACGGAGG	TAGAG M.len	rse
1019	ATTTAGGTGCAGCGT	TECETERTE	accacggage	TAGAG M.gas	tri
962	ATTTAGGTGCAGCGT	Tecerentic	accacggage	TAGAG M.kan	sasii
3408	ATTTAGGTGCAGCGT	Gechrefittd	TTGCCGGAGG	TAGAG M.sme	gmatis

		•	,		
	1050	1060	1070	1080	
2005	CAGCCAAACTCCG	AATGCCG-TGGT	G-TA-AAGC	GTGGCA	M.tuberculosis
1307	CAGCCAAACTCCG	AATGCCG-TGGT(G-TAMAAGC	GTGGCA 1	M.avium
1307	CAGCCAAACTCCG	AATGCCG-TGGT	G-TANAAGC	GTGGCA 1	M. naratubero
1401	CAGCCAAACTCCG	AATGCCGATAAG-	TGAAAGT	GTGGCA 1	M.nhlei
1323	CAGCCAAACTCCG	AATGCCG-TGGT	I-TANAAGC	GTGGCA 1	M.leprae
1098	CAGCCAAACTCCG	AATGCCG-TGGT(3-tahaFigc	GTGGCA 1	M. gastri
1041	CAGCCAAACTCCG	AATGCCG-TGGT	3-TAHA-GC	GTGGCA I	M.kansasii
3486	CAGCCAAACTCCG	aatgccggtaag(COALGAGI	goggaa i	M.smegmatis

Figure 1D

	1130	1140	1150	1160
2082	ACAGCCCAGATCGC	CGGCTAAGGCC	CCCAAGCGTG	GTGCTA M.tuberculos
1385	ACAGCCCAGATCGC	CGGCTAAGGCC	CCHAAGCGTG	TGCTA M. avium
1385	ACAGCCCAGATCGC	CGGCTAAGGCC	CCHAAGCGTG	TGCTA M. paratubero
1479	ACAGCCCAGATCGC	CGGCTAAGGCC	CCHAAGCGTG	TGCTA M.phlei
1401	ACAGCCCAGATCGC	CGGCTAAGGCC	CCHAAGCGTG	TGCTA M.lenrae
1175	ACAGCCCAGATCGC	CGGCTAAGGCC	CCHAAGCGTG	TGCTA M.gastri
1118	ACAGCCCAGATCGC	CGGCTAAGGCC	ССББББСССТС	TECTA M kongonii
3566	ACAGCCCAGATCGC	CGG[]TAAGGCC	CCHAAGCGT	TGTTA M. smegmatis

	1	290	13	00	13	10	13
2241	CTCAAGCA	CACCG	CCGAAGO	CGC	GGCACA'	CCF	CCTTGT-
544	CTCAAGCA						
44	CTCAAGCA						
538	CTCAAGCA						
60	CTCAAGCA					TICA	CCTTOTA
34	CTCAAGCA					7º	rccecb
77	CTCAAGCA						CCGCA
26	TCAAGCA	CACCG	CCGAAGC	CGC	GGAA	SCCA	ACGITTE
					·		
		.330	13	40	13	50	13
n		1					130
-	-GGTGGGT	Greec	PAGGGGA	GCG'	rcctc,	ATTO	AGCGAAG
3	-сстссет Ссстссят	erecer erecer	PAGGGGA PAGGGGA	GCG'	rccctc;	ATTC	AGCGAAG
33	-GGTGGGT GGGTGGAT GGGTGGAT	erecen erecen	PAGGGGA PAGGGGA	.GCG'	rccctc, rccccc,	ATTC	AGCGAAG AGCGAAG
33 33 76	- <u>GGTGGGT</u> CGGTGGAT CGGTGGT	Greeen eteeen eteeen	PAGGGGA PAGGGGA PAGGGGA	.GCG'	rcccrc; rccccc; rccccc;	ATTC ATTC ATTC	AGCGAAG AGCGAAG AGCGAAG
83 83 76	-GGTGGGT GGGTGGAT GGGTGGAT IGGCTGGAT GGGTGGAT	ereeer ereeer ereeer ereeer	PAGGGGA PAGGGGA PAGGGGA PAGGGGA	.GCG' .GCG' .GCG' .GCG'	recete, recece, recees, recete,	ATTC ATTC ATTC ATC	AGCGAAG AGCGAAG AGCGAAG EGIIGAAG
80 83 83 76 60 67	-GGTGGGT CGGTGGAT CGGTGGAT IGGCTGGAT GGGTGGAT	erecen erecen erecen erecen erecen	PAGGGGA PAGGGGA PAGGGGA PAGGGGA PAGGGGA	.GCG' .GCG' .GCG' .GCG'	TCCCTCA TCCCCCA TCCTCCA TCCTCA TCCTCA	ATTC ATTC ATTC ATC ATTC	AGCGAAG AGCGAAG AGCGAAG GGIGGAAG AGCGAAG
93 93 76 00	-GGTGGGT GGGTGGAT GGGTGGAT IGGCTGGAT GGGTGGAT	Green eteen eteen eteen eteen eteen	PAGGGA PAGGGGA PAGGGGA PAGGGGA PAGGGGA PAGGGGA	.GCG' .GCG' .GCG' .GCG' .GCG'	TCCCTCA TCCCCCA TCCCCCA TCCCTCA TCCCCCCCA TCCCCCCCA TCCCCCCCA TCCCCCCCA TCCCCCCCA TCCCCCCCA TCCCCCCCA TCCCCCCCA TCCCCCCCA TCCCCCCA TCCCCCCCA TCCCCCCCA TCCCCCCCA TCCCCCCCA TCCCCCCCA TCCCCCCCA TCCCCCCCA TCCCCCCCA TCCCCCCCA TCCCCCCCA TCCCCCCCA TCCCCCCA TCCCCCCA TCCCCCCA TCCCCCCA TCCCCCCA TCCCCCCA TCCCCCCA TCCCCCCA TCCCCCA TCCCCCCA TCCCCCA TCCCCCA TCCCCCA TCCCCCA TCCCCCA TCCCCCA TCCCCCA TCCCCCA TCCCCCA TCCCCCA TCCCCCA TCCCCCCA TCCCCCA TCCCCCCA TCCCCCA TCCCCCA TCCCCCA TCCCCCA TCCCCCA TCCCCCA TCCCCCA TCCCCCA TCCCCCA TCCCCCCA TCCCCCCA TCCCCCCA TCCCCCCA TCCCCCCA TCCCCCCA TCCCCCCA TCCCCCCA TCCCCCCA TCCCCCCA TCCCCCCA TCCCCCCA TCCCCCCA TCCCCCCA TCCCCCCCA TCCCCCCCA TCCCCCCCA TCCCCCCCA TCCCCCCCC	ATTO ATTO ATTO ATTO ATTO	AGCGAAG AGCGAAG AGCGAAG EGIIGAAG

Figure 1E

				
	1370	1380	1390	1400
2319	CCACCCCCCCCCCCC	CECCECCACCA		
1623	CACCEGGG GACCE	GTGGTGGAGGG	TGGGGGAGT	GAGAAT M.tuberculosis
	-1- 1	GTGGTGGAGGG	TGGGGGAGT	GAGAAT M.avium
1623	CL-cceeerevice	GTGGTGGAGGG	TGGGGGAGT	GAGAAT M.paratuberc.
1716	COGCCGAGTGAIICG	GTGGTGGAGGG	TGTGGGAGT	GAGAAT M.phlei
1640	CCTCCGGGTAACCG	GTGGTGGAGGG	TGGGGAAGT	GAGAAT M lenree
1402	CCGCCGGGTGACCG	GTGGTGGAGGF	TGGGGGAGT	GAGAAT M.gastri
1345	CTGCCGGGTGACCG	GTGGTGGAGG	TGGGGGAGT	GAGAAT M.kansasii
3796	COGCCGBGTATICEA	етестесь сес	TEMPERCA CONTRACTOR	GAGAAT M.smegmatis
		J0100100A000	TOFIGGGWGI	SAGAAT M.SMegmatis
		- · · ·	· · · · · · · · · · · · · · · · · · ·	· · ·
	1410	1420	1430	1440
onia			1430	1440
2359	GCAGGCATGAGTAG	CGACAAGGCAA	GTGAGAACC	TTGCCC M.tuberculosis
1662	GCAGGCATGAGTAG GCAGGCATGAGTAG	CGACAAGGCAA CGATAAGGCAA	GTGAGAACC'	TTGCCC M.tuberculosis
	GCAGGCATGAGTAG GCAGGCATGAGTAG GCAGGCATGAGTAG	CGACAAGGCAA CGATAAGGCAA CGATAAGGCAA	GTGAGAACC' GTGAGAACC' GTGAGAACC'	TTGCCC M.tuberculosis TTGCCC M.avium TTGCCC M.paratuberc
1662	GCAGGCATGAGTAG GCAGGCATGAGTAG GCAGGCATGAGTAG	CGACAAGGCAA CGATAAGGCAA CGATAAGGCAA	GTGAGAACC' GTGAGAACC' GTGAGAACC'	TTGCCC M.tuberculosis TTGCCC M.avium TTGCCC M.paratuberc
1662 1662	GCAGGCATGAGTAG GCAGGCATGAGTAG GCAGGCATGAGTAG GCAGGCATGAGTAG	CGACAAGGCAA CGATAAGGCAA CGATAAGGCAA CGATAAGGCAA	GTGAGAACC' GTGAGAACC' GTGAGAACC' GTGAGAACC'	TTGCCC M.tuberculosis TTGCCC M.avium TTGCCC M.paratuberc.
1662 1662 1756 1680	GCAGGCATGAGTAG GCAGGCATGAGTAG GCAGGCATGAGTAG GCAGGCATGAGTAG GCAGGCATGAGTAG	CGACAAGGCAA CGATAAGGCAA CGATAAGGCAA CGATAAGGCAA CGATAAGGCAA	GTGAGAACC' GTGAGAACC' GTGAGAACC' GTGAGAACC' GTGAGAACC'	TTGCCC M.tuberculosis FTGCCC M.avium FTGCCC M.paratuberc. FTTCCC M.phlei FTGCCC M.leprae
1662 1662 1756 1680 1442	GCAGGCATGAGTAG GCAGGCATGAGTAG GCAGGCATGAGTAG GCAGGCATGAGTAG GCAGGCATGAGTAG GCAGGCATGAGTAG	CGACAAGGCAA CGATAAGGCAA CGATAAGGCAA CGATAAGGCAA CGATAAGGCAA CGATAAGGCAA	GTGAGAACC' GTGAGAACC' GTGAGAACC' GTGAGAACC' GTGAGAACC' GTGAGAACC'	TTGCCC M.tuberculosis TTGCCC M.avium TTGCCC M.paratuberc. TTGCCC M.phlei TTGCCC M.leprae TTGCCC M.gastri
1662 1662 1756 1680 1442 1385	GCAGGCATGAGTAG GCAGGCATGAGTAG GCAGGCATGAGTAG GCAGGCATGAGTAG GCAGGCATGAGTAG GCAGGCATGAGTAG GCAGGCATGAGTAG	CGACAAGGCAA CGATAAGGCAA CGATAAGGCAA CGATAAGGCAA CGATAAGGCAA CGATAAGGCAA CGATAAGGCAA	GTGAGAACC' GTGAGAACC' GTGAGAACC' GTGAGAACC' GTGAGAACC' GTGAGAACC' GTGAGAACC'	TTGCCC M.tuberculosis TTGCCC M.avium TTGCCC M.paratuberc. TTTCCC M.phlei TTGCCC M.leprae TTGCCC M.gastri
1662 1662 1756 1680 1442	GCAGGCATGAGTAG GCAGGCATGAGTAG GCAGGCATGAGTAG GCAGGCATGAGTAG GCAGGCATGAGTAG GCAGGCATGAGTAG GCAGGCATGAGTAG	CGACAAGGCAA CGATAAGGCAA CGATAAGGCAA CGATAAGGCAA CGATAAGGCAA CGATAAGGCAA CGATAAGGCAA	GTGAGAACC' GTGAGAACC' GTGAGAACC' GTGAGAACC' GTGAGAACC' GTGAGAACC' GTGAGAACC'	TTGCCC M.tuberculosis TTGCCC M.avium TTGCCC M.paratuberc. TTGCCC M.phlei TTGCCC M.leprae TTGCCC M.gastri

	•	J			
	157		1590	1600	
2519	CCCCGTGAC	BAATCA-GCGGT	ACTAACCACCCAA	AACCG N	f.tuberculosis
1821	CGITCCOTGAIL	BAATCA-GCGGTI	ACTAACCACCCAA	AACCG N	1.avium
1821	CGICCOTGAIR	SAATCA-GCGGT	ACTAACCACCCAA	AACCG N	1.paratuberc.
1915	CGITCCOTGAIT	SAATCICATTON	CTAACCACCCAA	AACCM N	1 phlei
1840	CGCCCGTGAT	SAATCA-GCGGT	ACTOACCACCCAA	aaccg n	1.leprae
1602	CGCCCGTGAT	FAATCA-GCGGT	ACTAACCACCCAA	AACCG N	1.gastri
1545	CGCCCGTGATC	FAATCA-GCGGTI	ACTAACCACCCAA	AACCG M	1.kansasii
3996	CGLCCATCALC	Gaatca-gcggti	ACTAACCATICCAA	aacca m	1.smeqmatis

Figure 1F

	1610	1620	1630	1640
2558	GAT-CGATCAC-TC	CCCTTCGGGGG	TGTGGAGTT	C-TGG M.tuberculosis
1860	GAT-CGACCAT-TCC	CCTTCGGGG	C-GTGGCGDT	M-Dee M avium
1860	GAT-CGACCAIL-TCC	CCTTCGGGG	C-GTGGGGET	I-OGG M.paratuberc.
1955	GGG-CGATCE-ATC	FTTCGGGGF		TG-GG M phlai
1879	GAT-CGACCATATCO	CCTTCGGGGG	CTATEGAGGT	WEGG M lennes
1641	GAT-CGATCAC-TC	CCTTCGGGGG	A-GTGGAGGT	C-TCC M castri
1584	GAT-CGATCAC-TCC	CCTTCGGGGG	C-GTGGAGGT	C-TGG M.kansasii
4035	ACCGTGACCGCACCT	rrrceeeeE		GGTGG M.smegmatis
			, 101000011	police M. Smegmatis
				
	1650	1660	1670	1680
2594	GGCTGCGTGGGAACT	TCGCTGGTAG	TAGTCAAGCG	AAGGG M.tuberculosis
1896	GGCTGCGTGGGAQCT	TCGCTGGTAG	TAGTCAAGCA	AMGGG M.avium
1896	GGCTGCGTGGGACCT	TCGCTGGTAG	TAGTCAAGCA	AUGGG M. paratuherc
1986	GGCTGCGTGGGAQC	G-GTGGGTAG	TAGTCAAGCG	AUGGG M.phlei
1917	GGCTGCGTGGGAACT	TCGTTGGTAG	TAGTCAAGCG	AUGGG M.lenrae
1677	GGCTGCGTGGAGCT	TCGCTGGTAG	TAGTCAAGCG	AUGGG M. gastri
1620	GGCTGCGTGGAGCT	TCGCTGGTAG	TAGTCAAGCG	AIGGG M.kansasii
4071	GGCTGCATGGGACCT	TCGTTGGTAG	TAGTCAAGCG	AIGGG M.smegmatis
		U		- Elouebiicgiice 13
			γ	
	1690	1700	1710	1720
2634	-GTGACGCAGGAAGG	TAGCCGTACC	AGTCAGTGGT	AACA- M.tuberculosis
1936	-GTGACGCAGGAAGG	DAGCCGTACC	AGTCAGTGGT	AAMA- M.avium
1936	-GTGACGCAGGAAGG	CAGCCGTACC	AGTCAGTGGT	AATA- M.paratuberc.
2025	-GTGACGCAGGAAGG	TAGCCGTACC	AGTCAGTGGT	AATA- M.phlei
1957	-GTGACGCAGGAAGG	TAGCCGTACC	AGTCAGTGGT	AANA- M.leprae
1717	-GTGACGCAGGAAGG	DAGCCGTACC	AGTCAGTGGT	AATA- M.gastri
1660	-GTGACGCAGGAAGG	DAGCCGTACC	AGTCAGTGGT	AAMA- M.kansasii
4111	-GTGACGCAGGAAGG	TAGCCGTACC	GETCRETEET	AATA- M.smegmatis
	= = = = = = = = = = = = = = = = = = =		D-1041	with mismediatra

Figure 1G

	•	•			
	1730	1740	1750	1760	
1974 1974 2063 1995 1755 1698	-CTGGGGCAAGCC -CTGGGGCAAGCC -CTGGGGGAAACC -CTGGAGCAAGCC -CTGGGGCAAGCC	CGTAG - AGAG CGTAG - AGAG TGTAGGGGAGAG CGTAGGGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGA	CGATAGGCAAI CGATAGGCAAI GGATAGGCAAI CGATAGGCAAI CGATAGGCAAI	ATCCGT M.tub ATCCGT M.avi ATCCGT M.par ATCCGT M.phl ATCCGT M.lep ATCCGT M.gas	um atuberc. ei rae tri
4149	-cgeggiaacc	TGTAGGGAGTC	AGATAGGIAA	ATCCGT M.sme	gmatis

		970	1980	1990	200	<u> </u>
2908	AGGGGGACC	GGAATAT	GTGAACAC	CCTTGCGGTGG	SAGC	M.tuberculosis
2208	AGGGGGGCC	GGAATACC	GTGAACAC	CCTTGCGGTGG	SAGC	M.avium
2208	AGGGGGGCC	GGAATAC	GTGAACAC	CCTTGCGGTGG	AGC	M. paratuhero
2298	AGGGGGACC	Cacgtado	GTGAGGGC	TCTTGCGGGGG	AGC	M.phlei
2231	AGGGGGGCC	GGAATATC	GTGAACAC	CTTGCGGTGG	AGC	M.leprae
1910	•					M.gastri
1934	AGGGGGACC	GGAATACC	GTGAACAC	CCTTGCGGTGG	SAGC	M.kansasii
4385	AGGGGGACC	CACATEC	GTGTAAGC	CITTACGGCCC	AGC	M.smegmatis

		r			-		•
		10 .		20	2430	244	~
3345	ACCTCGACG	CCAGTT	GGGG	GGAGTC	TTGTTG	AAATACC	M.tuberculosis
284	ACCTCGACG	CCAGTT	GGGG	CGGAGTCG	TTGTTG	AAATACC	M.bovis
2645	GCACAGACG	CCAGTT	TGTG	GGAGTCG	TTGTTG	AAATACC	M.avium
393	ATACAGACG	CCAGTT	IGTA:	GGAGTCG	TTGTTG	AAATACC	M.intracellulare
2645	GCACAGACG	CCAGTT	IGIG	GGAGTCG	TTGTTG	AAATACC	M.paratuberc.
2737	GCTCGGACG	CCAGTT	dgggh	GGAGTCG	TTGTTG	AAATACC	M.phlei
2668	ACTTCGACG	ZIAGTT	GGG	GGAGTCG	TTGTTG	AAATACC	M.leprae
1910							M. gastri
2372	ACCTCAACG	CCAGTT	gggg	GGAGTCG	TTGTTG	AAATACC	M.kansasii
4822	GCTCACACG	CCAGTG	igggh	GGAGTCG	TTGTTG	AAATACC	M.smegmatis

Figure 1H

	24	50	2460	2470	248	0
3385	ACTCTGATC	GTATTGG	GCATCTAAC	CTCGAACCCT	CADTC	M.tuberculosis
324	ACTCTGATC	GTATTGG	GCATCTAAC	CTCGAACCCT	CABTC	M bovie
2685	ACTCTGATC	GTATTGG	ACAGCTAAC	GTCGAACCCT	-marc	M evium
433	ACTCTGATC	GTATTGG	ACACCTAAC	TCGAACCCT	-TIATC	M.intracellulare
2685	ACTCTGATC	STATTGG	ACACCTAAC	STCGAACCCT	-TATC	M. naratuhero
277.7	ACTCTGATC	STATTGG	GCCTCTAAC	CTCGGACCGT	THATC	M phlei
2708	ACTCTGATI	STATTGA	ACATCTAAC	CTCGAACCET	ATATC	M.lenrae
1910					-	M dastri
2412	ACTCTGATC	STATTGG	ACAGCTAAC	TCGAACCCT	מתמב	M kengacii
4862	ACTCTGATC	TATTGG	GOTTTAAC	TCGGACCGT	ATATC	M.smegmatis
			-	G 11		
		90	2500	2510	2520	
3425	GGGTTTAGG	SACAGTG	CCTGGCGGG	PAGTTTAACT	GGGC	M.tuberculosis
364	GGGTTTAGG	SACAGTG	CCTGGCGGG	PAGTTTAACT	REGEC	M. hovis
2724	GGGTTCACG	SACAGTG	CCTGGCGGG	PAGTTTAACT(GGGC	M.avium
472	GGGTTCACG	SACAGTG	CCTGGCGGG	PAGTTTAACT	GGGC	M.intracellulare
2724	GGGTTICIACGG	SACAGTG	CCTGGCGGG:	PAGTTTAACT	GGGC :	M.paratuberc.
2817	CGGTTCAGG(SACAGTG	CCTGGTGGG	PAGTTTAACT	GGGC	M.phlei
2748	GGTTTAGG	SACAGTG	cctggcggg	ragtttaact(GGGC 1	M.leprae
1910					1	M aggtri
2452	GGGTTDADGG	SACAGTG	CCTGGCGGGT	PAGTTTAACT	GGGC	M.kansasii
4902	DGGTTDAGGG	BACAGTG	ccreenees	TAGTTTAACTO	GGGC 1	M.smegmatis
	29	30 .	2940	2950	296	50
3864	AGTACGAGA	GACCGG	GACGGACGA	ACCTCTGGTG	CACCA	M.tuberculosis
3163	AGTACGAGA	GACCGG	GACGGACGA	ACCTCTGGT	TACCA	M. avium
3163	AGTACGAGA	GACCGG	GACGGACGA	ACCTCTGGT	TACCA	M.paratuberc.
3256	AGTACGAGA	GACCGG	GACGGACGA	ACCTCTGGTA	MACCA	M nhlei
3187	AGTACGAGA	GACCGG	GACGGACGA	ACCTCTGGT	TACCA	M.lenrae
1910						M.gastri
2891	AGTACGAGAG	GACCGG	GACGGACGA	ACCTCTAGTG	CACCA	M.kansasii
5342	AGTACGAGAG	GACCGG	GACGGACGA	ACCTCTGGTE	MACCA	M smeamatic

Figure 11

		2970	2980	2990	3000
3904	GTTGTCCC	GCAGGGG	ACCECTEGA	TAGCCACGTTO	GGT M.tuberculosis
3203	GTTGTCCC	accagggg	ACEGCTGGA	TAGCCACGTTC	GGA M. avium
3203	GTTGTCCC	ACCAGGGGC	acgctgga	TAGCCACGTTC	GGA M. paratibero
3296	GTTGTCCC	ACCAGGGGC	ACCGCTGGA	TAGCCACGTTC	GGA M.phlei
3227	GTTGTCIIC	MCCAGGGG	ACCGCTGGA	TAGCCACGTTC	GGA M.leprae
1910				_	M.gastri
2931	GTTGTCCC	ACCAGGGGC	ACCGCTGGA	TAGCHACGTTC	GGA M.kansasii
5382	GTTGTCCC	Mccaeeec	acegctega	TAGCCACGTTC	GGA M.smegmatis
			····		
	3	010	3020	3030	3040
3944	CAGGATAA	CCGCTGAAA	GCATCTAAG	CGGGAAACCTT	CTC M.tuberculosis
3243	CAGGATAA	CCGCTGAAA	GCATCTAAG	CGGGAAACCTT	CTC M. avium
3243	CAGGATAA	CCGCTGAAA	SCATCTAAG	CGGGAAACCTT	CTC M.paratuberc.
3336	CAGGATAAC	CCGCTGAAA	SCATCTAAG	CGGGAAACCTC	TTC Maphlei
3267	CAMGATAAC	CCCCTGAAA	SCATCTAAG	CGGGAAACCTT	CTC M.leprae
1910					M.gastri
2971	CAGGATAAC	CCGCTGAAA	SCATCTAAG (CGGGAAACCTT	CTC M.kansasii
5422	CAGGATAAC	CCGCTGAAA	SCATCTAAG (CGGGAAACCTC	TTC M.smegmatis
		•		· -	
	3	090	3100	3110	3120
4023	CCCGC-AGA	ACACGGGTT	CAATAGGTC	AGACCTGGAAG	CT M.tuberculosis
609	CCCGC-AGA	ACACGGGTT	CAATAGGTC	AGACCTGGAAG	CT M howis
3322	CCCGC-AGA	CACGGGAI	TGATAGGOC	AGACCTGGAAG	CT M avium
677	CCCGC-AGA	CACGGGTI	'CGATAGGIC	AGACCTGGBBG	CT M intradellulara
0022	CCCGC AGA	けんせつひゅつはげ	I GHI AGGAC	AGACCTGGAAG	CT M. paratuberc
3415	CCCGC-AGA	CACGGGA T	CGATAGACC	agacctg[a[g	CA M.phlei
3309		_			M.leprae
1910					M.gastri
3050	CCCGC-AGA	ACACGGGIT	'ggataggdc	agacctggaag	CT M.kansasii
5501	CCCGC-AGA	GCACGGGP I	TGATAGAC	AGACCTGGAAG	© M.smegmatis

Figure 1J

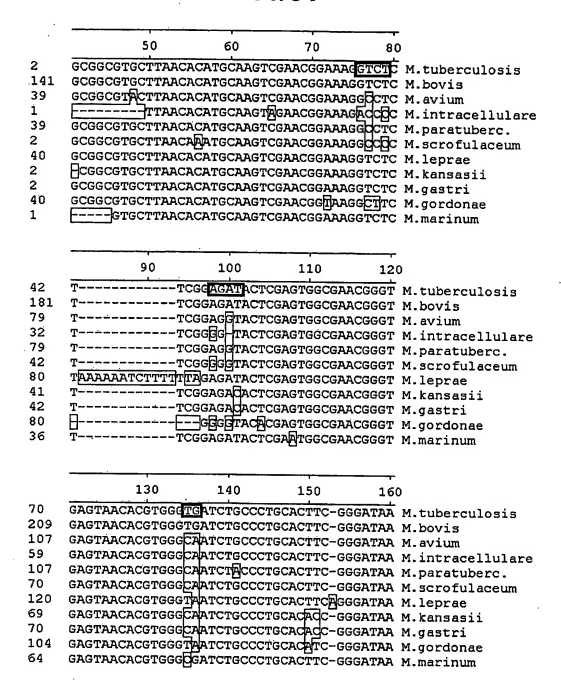


Figure 2A

	170	180	190	200
109	GCCTGGGAAACTG	GGTCTAATACCC	GATAGGACC	ACGGGA M. tuberculosis
248	GCCTGGGAAACTG	GGTCTAATACCG	GATAGGACC	ACGGGA M. hovis
146	GCCTGGGAAACTG	GGTCTAATACCG	GATAGGACO	CARCA M avium
98	GCCTGGGAAACTG	GGTCTAATACCE	GATAGGACC	TTAGG M.intracellulare
146	GCCTGGGAAACTG	GGTCTAATACCG	GATAGGACC	CAAGA M.paratuberc.
109	GCCTGGGAAACTG	GGTCTAATACCG	GATAGGACCI	CTTGG M scroful acoum
160	GCTTGGGAAACTG	GGTCTAATACCG	GATAGGACT	CAAGG M.leprae
108	GCCTGGGAAACTG	GGTCTAATACCG	GATAGGACC	CTTGG M.kansasii
109	GCCTGGGAAACTG	GGTCTAATACCG	GATAGGACC	CTIGG M.gastri
143	GCCTGGGAAACTG	GGTCTAATACCG	AATAGGACC	ACAGGA M.gordonae
103	GCCTGGGAAACTG	GGTCTAATACCG	GATAGGACC	ACGGGA M.marinum
				MILITAL THUM
	010	202		
	210	220	230	240
149	T GCATGTCTTGTG	GTGGAAAGCGCT	TTAGCGGTGT	GGGAT M.tuberculosis
288	TGCATGTCTTGTG	GTGGAAAGCGCT	TTAGCGGTGT	GGGAT M. hovis
186	GCATGTCTTGTG	GTGGAAAGC-TT	TT-ACGGTG1	GGGAT M.avium
138		CMCCSSSCO M		
186	DECATETETTE	GTGGAAAGC-TT	TT-GCGGTGT	GGGAT M.intracellulare AGAT M.paratuberc.
149	GCATGCTTGTG	GTGGAAAGCT	TTIGCGGTGT	GGGAT M.scrofulaceum
200			TTIGCGGTG	AGGAT M.leprae
148		このにこれれるとへし、一つ	mmmaaaaaaa	
149	GCATGCCTTGTG	STGGAAAGC-T	TINGCGGTGT	GGGAT M.kansasıı GGGAT M.gastri
183	CACATGTCCTATG	TGGAAAGC-TT	TT GCGGTGT	GGGAT M.gordonae
143	TICATGTCCTGTG	TGGAAAGF-CT	THICCGGTGT	GGGAT M.marinum
			- 150000101	GGGAT M.Malindm
	0.50	0.55		
	250	260	270	280
189	(AECCCGCGCCT)	ATCAGCTTGTTG	STGGGGTGAC	GGCCT M.tuberculosis
328	GAGCCCGCGGCCTA	\TCAGCTTGTTG(STGGGGTGAC	GGCCT M. bovis
224	GGGCCCGCGGCCT	ATCAGCTTGTTG(STGGGGTGAC	GGCCT M.avium
176	GGGCCCGCGGCCTZ	\TCAGCTTGTTG(STGGGGTGAM	GGCCT M.intracellulare
224	GEGCCCGCGCCTI	\TCAGCTTGTTG(STGGGGTGAC	GGCCT M paratuberd
187	GGCCCGCGCCT	TCAGCTAGTTG	STGGGGTG2M	GGCCT M.scrofulaceum
239	GEGCCCGCGCCTX	TCAGCTAATTA	STGGGGTAAC	GGCCT M.leprae
186	GGCCCGCGCCT	ATCAGCTTGTTG	₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽	GGCCT M.kansasii
187	GGCCCGCGCCT	ATCAGCTTGTTG(STGGGGGTGAC	GGCCT M destri
221	GE-CCCCCGCCT	ATCAGCTTGTTG(- 10000 годо.	GGCCT M.gastri
181	GGGCCGGCCTZ	TCAGCTTGTTG(പര്യപ്പോലാലെ	GGCCT M.marinum
			TOGGGTHYC	GGCCI M.Marinum

Figure 2B

	450	460	470	480
AAACC	TCTTTCACC	TCGACGAAGO	TCCGGGTTC	CTCGG M.tuberculos
AAACC	TCTTTCACC	TCGACGAAG	TCCGGGTTC:	TCTCGG M.bovis
AAACC	TCTTTCACC	TCGACGAAG	TCCGGGTTT	FCTCGG M.avium
AAACC	TCTTTCACC	TCGACGAAGG	TCCGGGTTH	CTCGG M.intracellu
AAACC	TCTTTCACC	TCGACGAAGG	TCCGGGTTT	TCTAGG M.paratubero
AAACC	TCTTTCACC	TCGACGAAGG	CTCACT	TTGTGG M.scrofulace
AAACC	TCTTTCACC	TCGACGAAGG	TCIGGGAAT	TCTCGG M.leprae
AAACC	TCTTTCACCE	TCGACGAAGG	TCCGGGTTC	CTCGG M.kansasii
AAACC	TCTTTCACCA	TCGACGAAGG	TCCGGGTTC	CTCGG M.gastri
AAACC	TCTTTCACCE	TCGACGAAGG	TCCGGGTTH.	CTCGG M.gordonee
AAACC	TCTTTCACCA	TCGACGAAGG	TICGGGTTH	CTCGG M.marinum
		~		
	1130	1140	1150	1160

			1100	110	•
1069	TCTCATGTTGCCAG	ACGTAATGGT	GGGGACTCG	TGAGAG	M.tuberculosis
1208	TCTCATGTTGCCAG	CACGTAATGGT	GGGGACTCG	TGAGAG	M. bovis
1104	TCTCATGTTGCCAG	GGTAATGCC	GGGGACTCG	TGAGAG	M.avium
1056	TCTCATGTTGCCAG	GGTAATGCC	GGGGACTCG	TGAGAG	M.intracellulare
1098	TCTCATGTTGCCAG	GGTAATGCA	GGGGACTCG	TGAGAG	M.paratuberc.
1064	TCTCATGTTGCCAG	GGGTAATGCC	GGGGACTCG	TGAGAG	M.scrofulaceum
1119	TCTCATGTTGCCAG	ACGTAATGGT	GGGGACTCG	TGAGAG	M.leprae
1066	TCTCATGTTGCCAG	GGGTAATGCC	GGGGACTCG	TGAGAG	M.kansasii
1067	TCTCATGTTGCCAG	GGGTAATGCC	GGGGACTCG	TGAGAG	M.gastri
1100	TCTCATGTTGCCAG	GGGTAATGCC	GGGGACTCG	TGAGAG	M.gordonae
1091	TCTCATGTTGCCAG	ACGTAATGGT	'GGGGACTCG'	TGAGAG	M.marinum

	12	50 12	260	1270	1280	
1189	CAATGGCCGG	TACAAAGGG	CTGCGATG	CCCCGAGGTT	AAG M.tuberculosi	S
1328	CAATGGCCGG	TACAAAGGG	CTGCGATG	CCGCGAGGTT	AAG M.bovis	
1224	CAATGGCCGG	TACAAAGGG	CTGCGATG	CCGTAAGGTTA	AAG M.avium	
1176	CAATGGCCGG	TACAAAGGG	CTGCGATG	CCGCAAGGTTA	AAG M.intracellul	are
1218	CAATGGCCGG	TACAAAGGG	CTGCGATG	ccg <u>r</u> aaggtta	AAG M.paratuberc.	
1184	CAATGGCCGG	TACAAAGGG	CTGCGATG	CCGCAAGGTTA	AG M.scrofulaceu	ım
1239	CAATGGCCGG	TACAAAGGG	CTGCGATG	CCGCAAGGTTA	AAG M.leprae	
					AAG M.kansasii	
1187		TACAAAGGG	CTGCGATG	CCGCGAGGTTA	VAG M.gastri	
1220	CAATGGCCGG	TACAAAGGG	CTGCGATG	CCGCGAGGTTA	NAG M.gordonae	
1181	CAATGGCCGG	TACAAAGGG	CTGCGATG	CCGCGAGGTTF	AAG M.marinum	

Figure 2C

					
	1290	1300	1310	1320	
1229	CGAATCCTTA-AAAC	CCGGTCTCAG	TTCGGATCG	GGGTCT M.tuberculosis	
1368	CGAATCCTTA-AAAG	CCGGTCTCAG	TTCGGATCG	GGGTCT M. boyis	
1264	CGAATCCTTTTAAAG	CCGGACTCAG	TTCGGATTG	GGGTCT M avium	
1216	CGAATCCTTTTAAAG	CCGGTCTCAG	TTCGGATTG	GGGTCT M.intracellula	ra
1258	CGAATCCTTTTAAAG	CCGGACTCAG	TTCGGATTG	GGGTCT M.paratuberc.	LC
1224	CGAATCCTTTTAAAG	CCGGTCTCAG	ттсесътсе	GGGTCT M.scrofulaceum	
1279	CGAATCCTTTTAAAG	CCGGTCTCAG	TTCGGATCG	GGGTCT M lenne	
1226	CGAATCCTTTTAAAG	CCGGTCTCAG	TTCGGATCG	GGGTCT M.kansasii	
1227	CGAATCCTTTTAAAG	CCGGTCTCAG	ТТСБСАТСС	GGTCT M destri	
1260	CGAATCCTTTTAAAG	CCGGTCTCAG	ТТСССТТССС	GGTCT M.gordonae	
1221	CGAATCCTTT-AAAG	CCGGTCTCAG	TTCCCATCC	GGTCT M.marinum	
		0000101010	TIOGGATCG	3661C1 M.Marinum	
	·				
		1340			
1268	GCAACTCGACCCCGI	GAAGTCGGAG	TCGCTAGTA	ATCGCA M.tuberculosis	
1407	GCAACTCGACCCCGT	GAAGTCGGAG	TCGCTAGTA	ATCGCA M.bovis	
1304	GCAACTCGACCCCAT	GAAGTCGGAG	TCGCTAGTA	ATCGCA M.avium	
1256	GCAACTCGACCCCAT	GAAGTCGGAG	TCGCTAGTA	ATCGCA M.intracellular	۹^
1298	GCAACTAGACCCAAT	GAAGTCGGAG	TCGCTAGTA	ATCGCA M.paratuberc.	
1264	GCAACTCGACCCCGT	GAAGTCGGAG	TCGCTAGTA	ATCGCA M.scrofulaceum	
1319	GCAACTCGACCCCGT	GAAGTCGGAG	TCGCTAGTA	ATCGCA M.lennee	
1266	GCAACTCGACCCCGT	GAAGTCGGAG	TCGCTAGTAR	ATCGCA M.kansasii	
1267	GCAACTCGACCCCGT	GAAGTCGGAG	TCGCTAGTAZ	ATCGCA M dastri	
1300	GCAACTCGACCCCGT	GAAGTCGGAG	тсестаетар	ATCGCA M.gordonae	
1260	GCAACTCGACCCCGT	GAAGTCGGAG	TCGCTAGTA	ATCGCA M.marinum	
				·- ·· · · · · · · · · · · · · · · · · ·	

Figure 2D

	50	60	70	80	
128 39 41 3559 5743	TECCGAACCCGGA TECCGAACCCGGA	agctaagcctgo agctaagcctgo agctaagcctgfi	cagogodga cagogodga loagogodga	TGATAC M.tuberc TGATAC M.bovis TGATAC M.phlei TGATAC M.leprae TGATAC M.smegma	
	90	100	110	120	
168 79 81 3599 5782	TGCCCCTCCGEG TGCCCTCACGGGG TGCCCATTCGGG-	TGGAAAAGT TGGAAAAGT	'agg <mark>g</mark> caccg 'aggacaccg	CCGAAC M.tuberc CCGAAC M.bovis CCGAAC M.phlei CCGAAC M.leprae	ulosis

Figure 3

•	90	100	110	120
382	GGGAGCTGTCAACCGA	GCATTGATO	CGAGGATTTC	CGAAT M.avium
382	GGGÄGCTGTCAACCGA	GCATTGATO	CGAGGATTTC	CGAAT M. paratuberc
1053	GGGAGCTGTCAACCGA	GCGTGGAT	CGAGGATTTC	CGAAT M. tuberculos
467	GGGAGCTGTCAACCGA	GCGTGGATC	CGAGGATTTC	CGAAT M.phlei
392	GGGAGCTGTCAACCGA	GCGTGGATC	CGAGGATTTC	CGAAT M.leprae
167	GGGAGCTGTCAACCGA	GCGTGGATC	CGAGGATTTC	CGAAT M.gastri
110	GGGAGCTGTCAACCGA	GCGTGGATC	CGAGGATTTC	CGAAT M.kansasii
2548	GGGAGCTGTCAACCGA	GCGTTGATC	CGAGGATGTC	CGAAT M. smegmatis

		7	, ,		
	170	180	190	200)
462	GAATATATAGGGTGCG	-GGAGGTA	ACCCCCCAAC	UCD 3 D	M. anadaam
	STATISTICS OF THE STATE OF THE	GGAGGIA	ACGCGGGGAAG:	TGAAA	M.avium
462	GAATATATAGGGTGCG	-ggaggta	ACGCGGGGAAG'	rgaaa	M. paratuberc
1133	GAATATATAGGGTGCG	-ggaggba	ACGCGGGGAAG'	TGAAA	M. tuberculogia
547	GAATATATAGGCGTTG	-ggggga	ACGCGGGGAAG'	IGAAA	M.phlei
472	GAATATATAGGGTTCG	-ggaggga	ACGCGGGGAAG'	IGAAA	M.leprae
247	GAATATATAGGGTGCG	-ggaggja	ACGCGGGGAAG!	IGAAA	M.gastri
190	GAATATATAGGGTGCG	-ggaggga	ACGCGGGGAAG'	IGAAA	M.kansasii
2628	GAATATATAGGCGTCT	-GGGGGA	acgcggggaag:	rgaaa	M.smegmatis

						
		250	. 260	270	280	þ
541	-GTCAGT	TAGTGGC	GAGCGAA	C-CGGAACA-G	GCTAAACCG	M.avium
541	-GTCAGT	PAGTGGC	GAGCGAA	C-CGGAACA-G	GCTAAACCG	M. paratubero
1212	-GCAAG1	ragtggc	GAGCGAAG	GCGGAACA-C	GCTAAACCG	M. tuberculosi
626	-GIGAGI	PAGTGGC	GAGCGAA	AGGGAGGATIC	GCTAAACCG	M.phlei
551	-GCAAGI	PAGTGGC	GAGCGAAG	GIGGAAIIATG	GCTAAACCG	M.leprae
326	-GTCAGI	AGTGGC	GAGCGAAG	GCGGAACATC	GCTAAACCG	M.gastri
269 2706	-GTAAGT	PAGTGGC	GAGCGAA	GCGGAACATG	GCTAAACCG	M.kansasii
2100	PP 1 PAG 1	AGTGGC	GAGCGAA(CGGAGGAIG	GCTAAACHG	M.smegmatis

Figure 4A

	· · · · · · · · · · · · · · · · · · ·	90	300	310	320	
578	CATG-CATG	GACAACCG	GGTAGGGGTT	TGTGTGCGGG	GT M.avium	
578	CATG-CATG	GACAACCG	GGTAGGGGTTC	- TOTOTOGO	GGT M.paratuberc.	
1250	CADG-CATG	SCC44FES	CGTACCCCTT(27GTGTGCGGG	GGT M.tuberculosis	
664	Care-Care		centrodoctro		GT M.phlei	
590	CALL CALC		201F0000110	TGTGTGCGG1		
365	CACA CATO	IT C THATCITA	GGTÄGGGGTT	erererecee[GT M.leprae	
	CACG-CATG	GGTGACCG	GGTAGGGGTT(FIGTGTGCGGG	GT M.gastri	
308	CADG-CATG	GEMAYCCG	GGTAGGGGTT	STGTGTGCGGG	GT M.kansasii	
2745	<u> </u>	TGATACCG	GGTAGGGGTT	TGTGTGCGGG	GT M.smegmatis	
					_	
		i	· · · · · · · · · · · · · · · · · · ·	-,		
	3	30	340	350	360	
617	TETEGGATT	CATATETE	TCACCTTCTACC	TITCCCITICT CO.	-GG M.avium	
617	TOTOGOTI	CAMARCAC	TCAGGICIACO	TEGGCTGAGG	-GG M.avium -GG M.paratuberc.	
1289	mcmcccr6	CATAIGIC	CAGCTCTACC	JIGGCIGAGG-	-GG M.paratuberc.	
	TGTGGGAG-	GATATGTC	TCAGCGCTACC	PGGCTGAGA-	GG M.tuberculosis	
703	TGTGGGGCC	Terene	CATICETCCE	Speedevieed	AG M.phlei	
629	TGTGGGATT	'GGTATGTC'	PCANCTCTAC(CTGGTTGAGG-	-GG M lenrae	
404	TGTGGGATC	GATACGTC	ICAGCTCTACC	CGGCTGAGG-	GG M.gastri	
347	TGTGGGATC	GATACGTC	PCAGCTCTAC (CGGCTGAGG-	GG M.kansasii	
2785	TGTGGGACC	TATOTHTC	POGGCTCTACC	TGGCTGFGAG	GG M.smegmatis	
	<u></u>		اس		goo :::::::::::::::::::::::::::::::::::	
		1	,	· · · · · · · · · · · · · · · · · · ·	·	
	3	70	380	390	400	
	•					
656	TAGTCAGAA	AGTGTCGT	ggttagcgga <i>i</i>	GTGGCCTGGG	AC M.avium	
656	TAGTCAGAA	agtgtcgt	GTTAGCGGAP	GTGGCCTGGG	AC M.paratuberc.	
1327	CAGTCAGAA	AGTGTCGT	egttagcgga <i>p</i>	GTGGCCTGGG	All M. tuberculosis	
742	TAGTGATAA	AGCAGTGT	gttaggtgap	GTGGCCTGGG	AI M.phlei	
668	TAGTCAGAA	AGTGCCGT	GTTAGCGGAP	ATGCCTCC	AI M.leprae	
443	CAGTCAGAA	AGTGTCGT	GTTADCGGA2		AI M.gastri	
386	CAGTCAGAA	AGTGTCGT		CUCCCCCCCC	AT M.kansasii	
2823	Charlet Char	701010010	COMPAGGGAN		M. Kansasii	
2023	Ava : Bugur	white Inc.	gttagcggaa	wriggchitege	AT M.smegmatis	

Figure 4B

		410	420	430	44	N
696	ecece	CCMACACCC	TCD CD CCCCC			-
696	GCCCCGC	CCULCACAC	GAGAGCCCGG	TACGCGAAA-	ACC	M.avium
1367	CCHCHC	CGTAGACGG	rgagageeegg	TACGCGAAA-	·ACC	M.paratuberc.
782	GGILCIGC	CGTAGACGG	rgagagcccgg	TACGCGAAA-	ACC	M. tuberculosis
708	GGITCHGC	CGTAGTGGG	GAGAGCCCGI	'AACHCGAAA-	ACA!	M.phlei
	GGCGIGC	CGTAGACGGT	GAGAGCCCAG	TACGCGAAA-	GCC	M.leprae
483	GGICIGC	CGTAGACGGT	GAGAGCCCGG	TACGIGAAA-	ACC	M.gastri
426	GRINOTICC	CGTAGACGGT	'GAGAGCCCGG	TACGMGAAA-	ACC	M kengacii
2863	eeccroc	CGTAGACGGT	rgagagcccgg	TACGIIGAAA-	ACC	M. smegmatis
		450	460	470	480	1
735	GGGG7 GG	ma a a a a a a a				=
735 735	CGGCACC	TGCCTTATAT	CAACA CCCGA	GTAGCAGCGG	GCC	M.avium
	CGGCACC	TGCCTTATAT	CAACACCCGA	GTAGCAGCGG	GCC	M. paratuberc.
820	Bolombook	TGCCTAGTAT	CAATTCCCGA	GTAGCAGCGG	GCC	M. tuberculosis
	IGCIGCO	Heciteticaca	GGTCCCGA	GTAGCAGCGG	GCC	M.phlei
747	TIGGCACC.	TGCCTTGTAT	CAATTCCCGA	GTAGCAGCGG	GCC	M.leprae
522	CGGCACC'	TGCCTTGTAT	CAATTCCCGA	GTAGCAGCGG	GCC	M.gastri
465	CGGCACC	TECCTTETAI	CAATTCCCGA	GTAGCAGCGG	GCC	M.kansasii
2902	CCACCIC.	TGIICTTGATG	GTGTTCCCGA	GTAGCAGCGG	GCC	M.smegmatis
			÷	-		
		520				
		570	580	590	600	
855	GAGGGAA!	TGGTGAAAAG	TACCCCGGGA	GGG-AGTGAA	ATA	M.avium
855	GAGGGAA:	TGGTGAAAAG	TACCCCGGGA	GGG-AGTGAA	מידמ	M naratubera
1526	GAGGGAA'	IGGTGAAAAG	TACCCCGGGA	ggggagtgaa:	AGA	M. tuberculosis
931	GAGGGAA!	IPGTGAAAAG	TACCCCGGGA	GGG-AGTGAA:	AGA	M.phlei
867	GAGGGAA!	IGGTGAAAAG	TACCCCGGGA	GGGGAGTGAA	ፈጥል	M lenrae
642	GAGGGAA	TGGTGAAAAG	TACCCCGGGA	ggggagtgaa:	AGA	Magatri
585	GAGGGAA1	IGGTGAAAAG	TACCCCGGGA	GGGGAGTGAA	AGA	M.kangasii
3022	GAGGGAAT	rggtgaaaag	TACCCCGGGA	GGGAGTGAA		M.smegmatis
					T	omediiar 12

Figure 4C

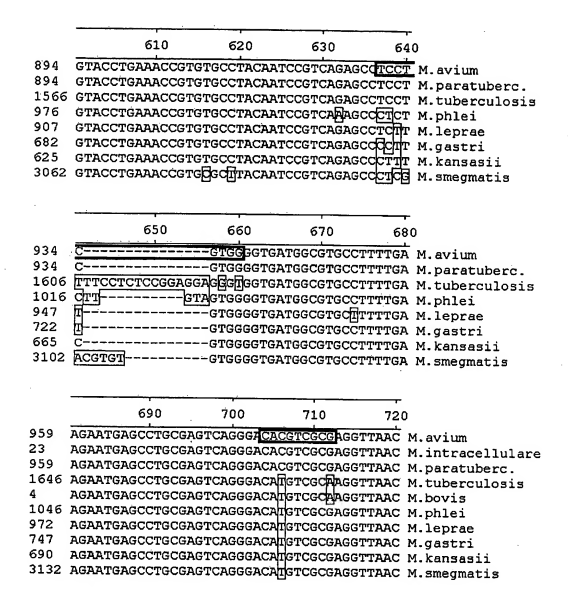


Figure 4D

	•	770	780	790	800	1
1039	CECATCCC	CTTTGGG		TETAGTGGCG'	rcr	M errium
103	CGCATCCC			700170000	ופו דפתי	M.intracellulare
	CGCATCCC			יים ואט וטטטטט ביים אניים מכרכי	ncm rgi	M.paratuberc.
1726	CGACCCAC	ACECE CATE	- حدددد سخمار	Manage Co	ran Igi	M.tuberculosis
84	CGACCCAC	CGCGCATA	CCCCCCTCTC	AATAGTGGCG'	IGI.	M. Cuberculosis
	CGTATCCA			TGTAGTGGTG	rer.	M.DOVIS
1052	CGMATCAC	FIGTGAGCG'	D	FTGTAGTGGCG:	I.G.I.	M.pniei
827		CGTAAGCG		TGTAGTGGCG'	rer.	M. leprae
770		CGCGAGCG		TIGIAGIGGCG	IGT.	m.gastri
3212		ACAAGAGT		TGTAGTGGCG	IGT	M. Kansasii
3212	COTINICOM	HCHHGHG10	31616	TGTAGTGGIIG	I'GT	M.smegmatis
						
	1	.050	1060	1070	108	0
1307	CAGCCAAA	CTCCGAATG	ссе-тесте	-TAAAAGCGTG	CCD	M evium
1307	CAGCCAAA	CTCCGAATG	CCG-TGGTG	-TANADACCATA	CCA	M.paratuberc.
2005	CAGCCAAA	CTCCGAATG	CCG-TGGTG	oroportuna -TaDaaccere	CCN	M.tuberculosis
1401	CAGCCAAA	СТССБЪЪТС	CCGDTDAG	-TGAAAGIGTG	CCN	M. cuberculosis
1323	CAGCCAAA	СТССБЪЪТС	CCG-TGGT	TAAAAGCGTG- TAAAAGCGTG	CCT	M. Johnson
1098	CAGCCAAA	СТССБЪТС	CCG-TGG1[]	-tafa-gcgtg	CCA	M.leprae
1041	CAGCCAAA	СТСССВВТС	CCG-MCGMG.	-maria-GCGTG	CCA	M.gastri M.kansasii
3486	CAGCCAAA	CTCCCAATG	CCGGTTTTTCC		GCA CED	M. kansasii M. smegmatis
3 100	CAGCCAAA	CICCGARIG	ccaalt err ac	-Charles Helle Cle	GAIA	M.smegmatis
						•
			~	·		•
		·				
	1	170	1180	1190	120	n
1425	ACTICO NA	N C C A ID C ID C	A CITICO CO A CIT			· -
1425	AGIGGAAA	ACCAMEMON	AGTCGCAGA	-GACAACCAGG	AGG	M.avium
2122	AGTCCGAAA	AGGATGTGT.	AGTCGCAGA	-gacaaccagg	AGG	M.paratuberc.
1510	VCIICODAY	AGGATGTGQ	AGTUGCAAA	-GACAACCAGG	AGG	M. tuberculosis
1019	AGTGGAAA	AGGATGTGO	AGTCGC GA	GACAACCAGG	AGG	M.phlei
1441	AGTGGAAA	AGGATGTGO	AGTCGCAMA-	-GACAACCAGG	AGG	M.leprae
1450	AGTGGGAA	AGGATGTGO	agtcgcaga.	-GACAACCAGG	AGG	M.gastri
TT28	AGTGGGAA	4GGATGTGC	AGTCGCAGA	GACAACCAGG	AGG	M.kansasii
3606	AGTGGAAA	aggatgtg <u>a</u>	AGTCGCAGA	ygapjaaccagg	AGG	M.smegmatis

Figure 4E

	1250	1260	1270	1280
1504	CTCACTGGTCAAGTG	ATTATGCGCC	GATAATGTAG	CGGGG M.avium
1504	CTCACTGGTCAAGTG	ATTATGCGCC	GATAATGTAG	CGGGG M. paratuberc.
2201	CTCACTGGTCAAGTG	ATTGTGCGCC	GATAATGTAG	CGGGG M.tuberculosis
1598	CTCACTGGTCAAGTG	attetececi	GATAATGTAG	CGGGG M.phlei
1520	CTCACTGGTCAAGTG	ATTGTGCGCC	GATAATGTAG	CGGGG M.leprae
1294		ATTGTGCGCC	GATAATGTAG	CGGGG M.gastri
1237	CTCACTGGTCAAGTG	ATTETECCC	GATAATGTAG	CGGGG M.kansasii
3686	TTCACTGGTCAAGTG	ATTGTGCGCC	GATATITGTEG	CGGGG M.smegmatis
		-		
			· · · · · · · · · · · · · · · · · · ·	
	1290	1300	1310	1320
1544	CTCAAGCACACCGCC	GAAGCCGCGG	CACATTCATC	TT-TA M.avium
1544	CTCAAGCACACCGCC	GAAGCCGCGG	CACATTCATC	TT-TA M.paratuberc.
2241	CTCAAGCACACCGCC	GAAGCCGCGG	CACATOCAOC	TTGT M. tuberculosis
1638	CTCAAGCACACCGCC	GAAGCCGCGG	CA-ATCAGO	Dring M.phlei
1560	CTCAAGCACACCGCC	GAAGCCGCGG	CACATTCACC	TTOTA M lenrae
1334	CTCAAGCACACCGCC	GAAGCCGCGA	CAFACC	FC-HA M dastri
1277	CTCAAGCACACCGCC	GAAGCCGCGA	CAACC	GC-A M.kansasii
3726	TTCAAGCACACCGCC	GAAGCCGCGG	BAL-GOCABC	
			Ft (arma w. swedwarts
		γ	 · · · · · · · · · · · · · · · ·	
	1330	1340	1350	1360
1583	CGGTGGATGTGGGTA	GGGAGCGTC	CCCCATTCAG	CGAAG M.avium
1583	CGGTGGATGTGGGTA	GGGAGCGTC	CCCCATTCAG	CGAAG M.paratuberc.
2280	FGGTGGGTGTGGGTA	GGGAGCGTC	CCTICATTCAG	CGAAG M.tuberculosis
1676	TGGCTGGTGTGGGTA	GGGAGCGTC	CITCCATCCCC	GAAG M.phlei
1600	GGTGGATGTGGGTA	GGGAGCGTH	CCICATTCAG	CGAAG M.leprae
1367	AGGTTGGGTA	GGGAGCGTC	CCICATTCAG	CGAAG M.gastri
1310				CGAAG M.kansasii
3764				GAAG M.smedmatis

Figure 4F

	-			
	1370	1380	1390	1400
1623	CT-CCGGGTGACCGG	TGGTGGAGG	TGGGGGAGTG	AGAAT M. avium
1623	CT-CCGGGTGATCGG	TGGTGGAGG	TGGGGGAGTG	AGAAT M.paratuberc.
2319	CEACCGGGTGACCGG	TGGTGGAGG	TGGGGGAGTG	AGAAT M. tuberculosis
1716	ccceActernices	TGGTGGAGG	TGTGGGAGTG	AGAAT M. nhlei
1640	CCTCCGGGTAACCGG	TGGTGGAGG	TGGGGDAGTG	AGAAT M lenree
1402	CCCCGGTGACCGG	TGGTGGAGG	TGGGGGAGTG	AGAAT M. gaatri
1345	CTGCCGGGTGACCGG	TGGTGGAGG	TGGGGGAGTG	AGAAT M. kansasii
3796	CCCCCACTATCCAC	TGGTGGAGG	TGTGGGAGTG	AGAAT M.smegmatis
				· · · · · · · · · · · · · · · · · · ·
			-	
				
	1530	1540	1550	1560
1781	CGATGGACAACGGGT	TGATATTCCC	GTACCCGTGT	ATGGG M.avium
1781	CGATGGACAACGGGT	TGATATTCCC	GTACCCGTGT	ATGGG M. naratuhera
2479	CGATGGACAACGGGT	TGATATTCCC	GTACCCGTGT	TGGG M.tuberculosis
1875	CGATGGACAACGGGT	TGATATTCCC	GTACCCGTGT	ATGAG M. phlei
1800	CGATGGACAACGGGT	TGATATTCCC	GTACCCGTGT	TGTG M. lenrae
1562	CGATGGACAACGGGT	TGATATTCCC	GTACCCGTGT	TGGG M.gastri
1505	CGATGGACAACGGGT	TGATATTCCC	GTACCCGTGT	TGGG M. kansasii
3956	CGATGGACAACGGGT	TGATATTCCC	GTACCCGTGT	ATG[[G M.smegmatis
				_
	1570	1580	1590	1600
1021			· -	
1021	CGTCCCTGATGAATC	A-GCGGTACT	AACCACCCAAI	ACCG M.avium
2510	CCCCCCTGATGAATC	A-GCGGTACT	AACCACCCAA	ACCG M.paratuberc.
1015	CCACCCACAGGAATC	RCD DECTACT	AACCACCCAAI	AACCG M.tuberculosis
1940	CGTCCTGATGAATC	TCATTOTECT	AACCACCCAAI	ACCI M.phlei
1602	CGCCCGTGATGAATC	n-ccccmrcm	PACCACCCAA	AACCG M.leprae
1.545	CGCCCTGATGAATC	n-ccccmrcm	AACCACCAA	AACCG M.gastri
300C	CGCCCTGATGAATC	A-GUGGTACT	AACCACCCAAI	AACCG M.kansasii
3330	CGICCHIGATGAATC	4-GCGGTACT	AACCAIICCAAI	AACCA M. smegmatis

Figure 4G

	1610	1620	1630	1640
1860	GAT-CGACCAT-TCC	CCTTCCCCC	C-CECCCC	
1860	GAT-CGACCAT-TCC	CC11CGGGG	C-GIGGCGA	TT-CGG M.avium TT-CGG M.paratuberc.
2558	GAT-CGARCAC-TCC	70011000000 70011000000		mc-mcc w tubercation
1955	GEG-CGANCE-ATTCC	EDTTCGGGG		TG-TGG M.tuberculosis TTG-GG M.phlei
1879	GAT-CGACCATATCC		CENT COLC	TIB-GG M. DILEI
1641	CAT CONCONTENCO	CCTTCGGGG	D Charle	TT-CGG M. Leprae
1504	GAT CGATCAC TCC		A-GIGGAGG	TO-TGG M.gastri TO-TGG M.kansasii
4035	PCCCUCACCCO CC		C-GTGGAGG	TO-1166 M. Kansasii
4033	WCCQ IQUCCOCHOCI		-Jenedocell	TGGTGG M.smegmatis
				
	1650	1660	1670	1680
1896	GGCTGCGTGGGACCT	TCGCTGGTAG	TAGTCAAGO	AATEGG M.avium
1896	GGCTGCGTGGGACCT	TCGCTGGTAG	TAGTCAAGC	AATGGG M Daratuberc
2594	GGCTGCGTGGGAACT	TCGCTGGTAG	TAGTCAAGO	GAAGGG M.tuberculosis
1986	GGCTGCGTGGGACCC	G-GIGGGTAG	TAGTCAAGC	GATEGE M phlei
1917	GGCTGCGTGGGAACT	TCGTTGGTAG	TAGTCAAGC	GATGGG M.lenrae
1677	GCTGCGTGGAACT GGCTGCGTGGAGCCT	TCGCTGGTAG	TAGTCAAGC	GATGGG M.gastri
1620	GGCTGCGTGGAGCCT	TCGCTGGTAG	TAGTCAAGC	GATGGG M.kansasii
4071	GGCTGCATGGGACCT	TCGTTGGTAG	TAGTCAAGC	GATGGG M.smegmatis
				3.1000 11.5mcgmat13
	•			
	1690	1700	1710	1720
1936	-GTGACGCAGGAAGG	CAGCCGTACC	AGTCAGTGG'	TAATA- M.avium
1936	-GTGACGCAGGAAGG	CAGCCGTACC	AGTCAGTGG'	TAATA- M.avium
1936	-GTGACGCAGGAAGG	CAGCCGTACC	AGTCAGTGG'	TAATA- M.avium
1936 2634	-GTGACGCAGGAAGG -GTGACGCAGGAAGG -GTGACGCAGGAAGG	CAGCCGTACC CAGCCGTACC	AGTCAGTGG' AGTCAGTGG'	TAATA- M.avium TAATA- M.paratuberc. TAADA- M.tuberculosis
1936 2634 2025	-GTGACGCAGGAAGG -GTGACGCAGGAAGG -GTGACGCAGGAAGG -GTGACGCAGGAAGG	CAGCCGTACC CAGCCGTACC TAGCCGTACC	AGTCAGTGG' AGTCAGTGG' AGTCAGTGG' AGTCAGTGG'	TAATA- M.avium TAATA- M.paratuberc. TAADA- M.tuberculosis
1936 2634 2025 1957	-GTGACGCAGGAAGG -GTGACGCAGGAAGG -GTGACGCAGGAAGG -GTGACGCAGGAAGG	CAGCCGTACO CAGCCGTACO TAGCCGTACO TAGCCGTACO TAGCCGTACO	AGTCAGTGG' AGTCAGTGG' AGTCAGTGG' AGTCAGTGG' AGTCAGTGG'	TAATA- M.avium TAATA- M.paratuberc. TAAGA- M.tuberculosis TAATA- M.phlei TAATA- M.leprae
1936 2634 2025 1957 1717	-GTGACGCAGGAAGG -GTGACGCAGGAAGG -GTGACGCAGGAAGG -GTGACGCAGGAAGG -GTGACGCAGGAAGG	CAGCCGTACO CAGCCGTACO TAGCCGTACO TAGCCGTACO TAGCCGTACO CAGCCGTACO	AGTCAGTGG' AGTCAGTGG' AGTCAGTGG' AGTCAGTGG' AGTCAGTGG'	TAATA- M.avium TAATA- M.paratuberc. TAAGA- M.tuberculosis TAATA- M.phlei TAATA- M.leprae TAATA- M.gastri
1936 2634 2025 1957 1717 1660	-GTGACGCAGGAAGG -GTGACGCAGGAAGG -GTGACGCAGGAAGG -GTGACGCAGGAAGG -GTGACGCAGGAAGG -GTGACGCAGGAAGG	CAGCCGTACO CAGCCGTACO TAGCCGTACO TAGCCGTACO CAGCCGTACO CAGCCGTACO	AGTCAGTGG' AGTCAGTGG' AGTCAGTGG' AGTCAGTGG' AGTCAGTGG' AGTCAGTGG' AGTCAGTGG'	TAATA- M.avium TAATA- M.paratuberc. TAADA- M.tuberculosis TAATA- M.phlei TAATA- M.leprae TAATA- M.gastri TAATA- M.kansasii
1936 2634 2025 1957 1717 1660	-GTGACGCAGGAAGG -GTGACGCAGGAAGG -GTGACGCAGGAAGG -GTGACGCAGGAAGG -GTGACGCAGGAAGG -GTGACGCAGGAAGG	CAGCCGTACO CAGCCGTACO TAGCCGTACO TAGCCGTACO CAGCCGTACO CAGCCGTACO	AGTCAGTGG' AGTCAGTGG' AGTCAGTGG' AGTCAGTGG' AGTCAGTGG' AGTCAGTGG' AGTCAGTGG'	TAATA- M.avium TAATA- M.paratuberc. TAAGA- M.tuberculosis TAATA- M.phlei TAATA- M.leprae TAATA- M.gastri
1936 2634 2025 1957 1717 1660	-GTGACGCAGGAAGG -GTGACGCAGGAAGG -GTGACGCAGGAAGG -GTGACGCAGGAAGG -GTGACGCAGGAAGG -GTGACGCAGGAAGG -GTGACGCAGGAAGG	CAGCCGTACO CAGCCGTACO TAGCCGTACO TAGCCGTACO CAGCCGTACO CAGCCGTACO	AGTCAGTGG' AGTCAGTGG' AGTCAGTGG' AGTCAGTGG' AGTCAGTGG' AGTCAGTGG' AGTCAGTGG'	TAATA- M.avium TAATA- M.paratuberc. TAADA- M.tuberculosis TAATA- M.phlei TAATA- M.leprae TAATA- M.gastri TAATA- M.kansasii TAATA- M.smegmatis
1936 2634 2025 1957 1717 1660 4111	-GTGACGCAGGAAGG -GTGACGCAGGAAGG -GTGACGCAGGAAGG -GTGACGCAGGAAGG -GTGACGCAGGAAGG -GTGACGCAGGAAGG -GTGACGCAGGAAGG -GTGACGCAGGAAGG	CAGCCGTACC CAGCCGTACC TAGCCGTACC TAGCCGTACC CAGCCGTACC CAGCCGTACC TAGCCGTACC	AGTCAGTGG' AGTCAGTGG' AGTCAGTGG' AGTCAGTGG' AGTCAGTGG' AGTCAGTGG' AGTCAGTGG' AGTCAGTGG'	TAATA- M.avium TAATA- M.paratuberc. TAADA- M.tuberculosis TAATA- M.phlei TAATA- M.leprae TAATA- M.gastri TAATA- M.kansasii TAATA- M.smegmatis
1936 2634 2025 1957 1717 1660 4111	-GTGACGCAGGAAGG -GTGACGCAGGAAGG -GTGACGCAGGAAGG -GTGACGCAGGAAGG -GTGACGCAGGAAGG -GTGACGCAGGAAGG -GTGACGCAGGAAGG -GTGACGCAGGAAGG -GTGACGCAGGAAGG	CAGCCGTACC CAGCCGTACC TAGCCGTACC TAGCCGTACC CAGCCGTACC TAGCCGTACC TAGCCGTACC	AGTCAGTGG' AGTCAGTGG' AGTCAGTGG' AGTCAGTGG' AGTCAGTGG' AGTCAGTGG' AGTCAGTGG' AGTCAGTGG'	TAATA- M.avium TAATA- M.paratuberc. TAADA- M.tuberculosis TAATA- M.phlei TAATA- M.leprae TAATA- M.gastri TAATA- M.kansasii TAATA- M.smegmatis
1936 2634 2025 1957 1717 1660 4111	-GTGACGCAGGAAGG	CAGCCGTACC CAGCCGTACC TAGCCGTACC TAGCCGTACC CAGCCGTACC TAGCCGTACC TAGCCGTACC TAGCCGTACC	AGTCAGTGG'	TAATA- M.avium TAATA- M.paratuberc. TAADA- M.tuberculosis TAATA- M.phlei TAATA- M.leprae TAATA- M.gastri TAATA- M.kansasii TAATA- M.smegmatis 1760 ATCCGT M.avium
1936 2634 2025 1957 1717 1660 4111 1974 1974 2672	-GTGACGCAGGAAGG -GTGACGCAGGAAGG -GTGACGCAGGAAGG -GTGACGCAGGAAGG -GTGACGCAGGAAGG -GTGACGCAGGAAGG -GTGACGCAGGAAGG -GTGACGCAGGAAGG -GTGACGCAGGAAGG -CTGGGGCAAGCCCG	CAGCCGTACC CAGCCGTACC TAGCCGTACC TAGCCGTACC CAGCCGTACC TAGCCGTACC TAGCCGTACC TAGCCGTACC TAGCCGTACC TAGCCGTACC TAGCCGTACC	AGTCAGTGG'	TAATA- M.avium TAATA- M.paratuberc. TAAGA- M.tuberculosis TAATA- M.phlei TAATA- M.leprae TAATA- M.gastri TAATA- M.kansasii TAATA- M.smegmatis 1760 ATCCGT M.avium ATCCGT M.paratuberc. ATCCGT M.tuberculosis
1936 2634 2025 1957 1717 1660 4111 1974 1974 2672 2063	-GTGACGCAGGAAGG -GTGACGCAGGAAGG -GTGACGCAGGAAGG -GTGACGCAGGAAGG -GTGACGCAGGAAGG -GTGACGCAGGAAGG -GTGACGCAGGAAGG -GTGACGCAGGAAGG -GTGACGCAGGAAGG -CTGGGGCAAGCCCG -CTGGGGCAAGCCCG	CAGCCGTACC CAGCCGTACC TAGCCGTACC TAGCCGTACC CAGCCGTACC TAGCCGTACC TAGCCGTACC TAGCCGTACC TAGCCGTACC TAGCCGTACC TAGCCGTACC TAGCCGTACC TAGCCGTACC	AGTCAGTGG' AGTCAGTAGGCAAI	TAATA- M.avium TAATA- M.paratuberc. TAAGA- M.tuberculosis TAATA- M.phlei TAATA- M.leprae TAATA- M.gastri TAATA- M.kansasii TAATA- M.smegmatis 1760 ATCCGT M.avium ATCCGT M.paratuberc. ATCCGT M.tuberculosis
1936 2634 2025 1957 1717 1660 4111 1974 1974 2672 2063 1995	-GTGACGCAGGAAGG -GTGACGCAGGAAGG -GTGACGCAGGAAGG -GTGACGCAGGAAGG -GTGACGCAGGAAGG -GTGACGCAGGAAGG -GTGACGCAGGAAGG -GTGACGCAGGAAGG -GTGACGCAGGAAGG -CTGGGGCAAGCCCG -CTGGGGCAAGCCCG -CTGGGGCAAGCCGG -CTGGGGCTAARCCTG	CAGCCGTACC CAGCCGTACC TAGCCGTACC TAGCCGTACC CAGCCGTACC CAGCCGTACC TAGCCGTACC TAGCCGTACC TAGCCGTACC TAGCCGTACC TAGCCGTACC TAGCCGTACC TAGCCGTACC TAGCCGTACC TAGCCGTACC	AGTCAGTGG' AGTCAGTAGGCAAI GATAGGCAAI	TAATA- M.avium TAATA- M.paratuberc. TAAGA- M.tuberculosis TAATA- M.phlei TAATA- M.leprae TAATA- M.gastri TAATA- M.kansasii TAATA- M.smegmatis 1760 ATCCGT M.avium ATCCGT M.paratuberc. ATCCGT M.tuberculosis ATCCGT M.phlei
1936 2634 2025 1957 1717 1660 4111 1974 1974 2672 2063 1995 1755	-GTGACGCAGGAAGG -GTGACGCAGGAAGG -GTGACGCAGGAAGG -GTGACGCAGGAAGG -GTGACGCAGGAAGG -GTGACGCAGGAAGG -GTGACGCAGGAAGG -GTGACGCAGGAAGG -GTGACGCAGGAAGG -CTGGGGCAAGCCCG -CTGGGGCAAGCCCG -CTGGGGCAAGCCCG -CTGGGGCAAGCCCG -CTGGGGCAAGCCCG	CAGCCGTACC CAGCCGTACC TAGCCGTACC TAGCCGTACC CAGCCGTACC TAGCCGTACC TAGCCGTACC TAGCCGTACC TAGCCGTACC TAGCCGTACC TAGCCGTACC TAGCCGTACC TAGCCGTACC TAGCCGTACC TAGCCGAGAGC TAGGCAGAGAGC TAGGCAGAGAGC TAGGCAGAGAGC TAGGCAGAGAGC	AGTCAGTGG' AGTCAGGCAAI GATAGGCAAI GATAGGCAAI	TAATA- M.avium TAATA- M.paratuberc. TAADA- M.tuberculosis TAATA- M.phlei TAATA- M.leprae TAATA- M.gastri TAATA- M.kansasii TAATA- M.smegmatis 1760 ATCCGT M.avium ATCCGT M.paratuberc. ATCCGT M.tuberculosis ATCCGT M.phlei ATCCGT M.leprae ATCCGT M.gastri
1936 2634 2025 1957 1717 1660 4111 1974 1974 2672 2063 1995 1755 1698	-GTGACGCAGGAAGG -GTGACGCAGGAAGG -GTGACGCAGGAAGG -GTGACGCAGGAAGG -GTGACGCAGGAAGG -GTGACGCAGGAAGG -GTGACGCAGGAAGG -GTGACGCAGGAAGG -GTGACGCAGGAAGG -CTGGGGCAAGCCCG -CTGGGGCAAGCCCG -CTGGGGCAAGCCGG -CTGGGGCAAGCCGG -CTGGGGCAAGCCAG	CAGCCGTACC CAGCCGTACC TAGCCGTACC TAGCCGTACC CAGCCGTACC TAGCCGTACC TAGCCGTACC TAGCCGTACC TAGCCGTACC TAGCCGTACC TAGCGAGAGC TAGGGAGAGC TAGGGAGAGAGC TAGGGAGAGAGC TAGGGAGAGAGC TAGGGAGAGAGC TAGGGAGAGAGC TAGGGAGAGAGC TAGGGAGAGAG	AGTCAGTGG' AGTAGGCAAAAAGGAAAAGGAAAAGGAAAAGGAAAAAGAAAAAA	TAATA- M.avium TAATA- M.paratuberc. TAAGA- M.tuberculosis TAATA- M.phlei TAATA- M.leprae TAATA- M.gastri TAATA- M.kansasii TAATA- M.smegmatis 1760 ATCCGT M.avium ATCCGT M.paratuberc. ATCCGT M.tuberculosis ATCCGT M.phlei

Figure 4H

```
1810
                           1820
                                      1830
                                                 1840
 2051 CG-AATTCGGTGATCCTCTGCTGCCAAGAAAGCCTCTA- M.avium
 2051 CG-AATTCGGTGATCCTCTGCTGCCAAGAAAGCCTCTA- M.paratuberc.
 2751 CG-AATTCGGTGATCCTCTGCTGCCAAGAAAGCCTCTA- M.tuberculosis
 2141 CG-AATTCGGTGATCCTATGCTGTCGAGAAAAGCCTCTA- M.phlei
2074 CG-AATTCGGTAAGCCTCTGCTGCCAAGAAAGCCTCTA- M.leprae
 1834 CG-AATTCGGTGATCCTCTGCTGCCAAGAAAGCCTCTA- M.gastri
1777 CG-AATTCGGTGATCCTCTGCTGCCAAGAAAAGCCTCTA- M.kansasii
 4228 CG-AATTCGGTGATCCTATGCTGCCGAGAAAAGCCTCTA- M.smegmatis
               1850
                          1860
                                     1870
                                                 1880
2089 GCGAGCACATACACGGCCCGTACCCCAAACCAACACAGGT M.avium
2089 GCGAGCACATACACTGCCCGTACCCCAAACCAACACAGGT M.paratuberc.
2789 GCGAGCACACGCCCGTACCCCAAACCGACACGGT M.tuberculosis
2179 GCAAGCGCATACACGGCCCGTACCCCAAACCAACACAGGT M.phlei
2112 GCGAGCATACATGCGGCCCGTACCCCAAACCGACACAGGT M.leprae
1872 GCGAGCACACACGGCCCGTACCCCAAACCGACACAGG M.gastri
1815 GCGAGCACACACGGCCCGTACCCCAAACCGACACAGGT M.kansasii
4266 GCGAGGACATACACGGCCCGTACCCCAAACCAACACAGGT M.smegmatis
              1970
                          1980
                                     1990
                                                2000
2208 AGGGGCCCGGAATACCGTGAACACCCTTGCGGTGGGAGC M.avium
2208 AGGGGGCCCGGAATACCGTGAACACCCTTGCGGTGGGAGC M.paratuberc.
2908 AGGGGGACCGGAATATCGTGAACACCCTTGCGGTGGGAGC M.tuberculosis
2298 AGGGGGACCCACGTACCGTGAGGCCTTTGCGGGGGGAGC M.phlei
2231 AGGGGGCCGGAATATCGTGAACACCCTTGCGGTGGGAGC M.leprae
1910
                                                    M.gastri
1934 AGGGGGACCGGAATACCGTGAACACCCTTGCGGTGGGAGC M.kansasii
4385 AGGGGGACCCACATGGCGTGTAAGCCTTTACGGCCCAAGC M.smegmatis
              2010
                          2020
                                     2030
                                                2040
2248 GGGATTCGGCCCGCAGAAACCAGTGGGTAGCGACT-GTTTA M. avium
2248 GGGATTCGGCCGCAGAAACCAGTGGGTAGCGACT-GTTTA M.paratuberc.
2948 GGGAT CGGTCGCAGAAACCAGTGAGGAGCGACT-GTTTA M.tuberculosis
2338 GGGGTGGCAGAAACCAGTGAGGAGCGACT-GTTTA M.phlei
2271 GGGAT CGGTCGCAGAGACCAGTGAGAGCGACT-GTTTA M.leprae
1974 GGGATTCGGTCGCAGAAACCAGTGAGAGCGACTTGTTTA M.kansasii
4425 GTGAGTGGCAGAAACCAGTGAGAGCGACT-GTTTA M.smegmatis
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Figure 41

	2130	2140	2150	2160
2367	CCGTTAACCCGT -	AAGGGTGAAGG	CGGAGAATTT	AAGCCC M.avium
2367	CCGTTAACCCGT	AAGGGTGAAG (CGGAGAATTT	AAGCCC M.paratuberc.
3067	CCGTTAACCCGQ:	AAGGGTGAAG(CGGAGAATTT	AAGCCC M.tuberculosis
2457	CCGTTAACCCHTTC	ggggtgaag(GGAGAATTT	AAGCCC M nhlei
2390	CIGTTAACCCGA	AAGGGTGAAGG	GGAGAATTT	AAGCCC M.leprae
1910				M destri
2094	CCGTTAACCCGQ	AAGGGTGAAGC	GGAGAATTT	ABGCCC M kangagii
4544	CCGTTAACCCCCTT	GGGTGAAG	GGAGAATTT	AAGCCC M.smegmatis
	· · · · · · · · · · · · · · · · · · ·			-10000 Dineginating
			•	
				
	2250	2260	2270	2280
2485	GTAACGACTTCCCA	CTGTCTCAAC	CATAGACTO	GCGAA M.avium
2485	GTAACGACTTCCCA	ACTGTCTCAAC	CATAGACTC	GCGAA M. paratuhero
3185	GTAACGACTTCTCAI	CTGTCTCAAC	CATAGACTO	GCGAA M.tuberculosis
2577	GTAACGACTTCICAL	CTGTCTCAAC	CATAGACTC	GCGAA M.phlei
2508	GTAACGACTTCTCAL	CTGTCTCAAC	CATAGACTC	GCGAA M.lenrae
1910	•			M.gestri
2212	GTAACGACTTCTCAA	CTGTCTCAAC	CATAGACTC	GCGAA M.kansasii
4663	GTAACGACTTCTCAA	CTGTCTCAAC	HATAGACTCO	GCGAA M.smegmatis
	_			
	2370	2380	2390	2400
2605	GTTCGGTACGGTTTG	TGTAGGATAG	GTGGGAGACT	TTGAA M.avium
2605	GTTCGGTACGGTTTG	TGTAGGATAG	GTGGGAGAC1	TTGAA M.paratuberc.
3305	GTTCGGTACGGTTTG	TGTAGGATAG	GTGGGAGACT	GTGAA M.tuberculosis
2697	GOTCGATACGGTTTG	TGTAGGATAG	GTGGGAGACT	GTGAA M.phlei
2628	GTTCGGTGCGGTTTG	TGTAGGATAG	GTGGGAGACT	GTGAA M.leprae

Figure 4J

2332 GTTCGGTACGGTTTGTGTAGGATAGGTGGGAGACTGTGAA M.kansasii 4782 GGTCGATACGGTTGTAGGATAGGTGGGAGACTGTGAA M.smegmatis

				
	2410	2420	2430	2440
2645	GCACAFACGCCAGTT	TETETEGAG	ГССТТСТТСЬВ	ATACC M avium
393	ATACAGACGCCAGTT	'TGTATGGAG'	ϹႺͲͲ;;ͲϯϦϹϽϽ	ATACC M intracellulare
2645	GCACAGACGCCAGTT	'TGTGTGGAG'	AADTTGTTGD1	ATACC M. paratuberc
3345	ACCTOGACGCCAGTT	GGGGGGGAGT	CGTTGTTGAA	ATACC M.tuberculosis
284	ACCTOGACGCCAGTT	GGGGGGGAG'	CGTTGTTGAA	ATACC M.bovis
2737	GCICGGACGCCAGTT	DGGGTGGAGT	ГССТТСТТСЭ	ATACC M phlei
2668	ACTTCGACGCHAGTT	GGGGTGGAG	AAƏTTƏTTƏƏ	ATACC M.leprae
1910				M costni
2372	ACCTCAACGCCAGTT	GGGGTGGAGT	rcgttgttgaa	ATACC M.kansasii
4822	GCIICACACGCCAGTE	TGGGTGGAGT	CGTTGTTGAA	ATACC M.kansasii ATACC M.smegmatis
	0.450			
•	2450	2460	2470	2480
2685	ACTCTGATCGTATTG	GACACCTAA	CGTCGAACCCT	-TATC M.avium
433	ACTCTGATCGTATTG	GACACCTAAC	GTCGAACCCT	-TATC M.intracellulare
2685	ACTCTGATCGTATTG	GACACCTAAC	GTCGAACCCT	-TATC M.paratuberc.
3385	ACTCTGATCGTATTG	GGCAIICTAAC	CTCGAACCCT	GAATC M.tuberculosis
324	ACTCTGATCGTATTG	GECATICTAAC	COTCGAACCCT	GAATC M.bovis
2777	ACTCTGATCGTATTG	GGCCTCTAAC	CTCGGACCGT	GGATC M.phlei
2708	ACTCTGATTGTATTG	PACALICTAAC	CTCGAACCGT	ATATC M.leprae
TATO				M. dastri
2412	ACTCTGATCGTATTG	GACACCTAAC	GTCGAACCCT	GAATC M.kansasii
4862	ACTCTGATCGTATTG	GGCCTCTAAC	CTCGGACCGT	MTATC M. smegmatis
		_		
	2690	0700	0.710	
		2700	2710	2720
2924	GGTGTCACTCAACGG	ATAAAAGGT.	ACCCCGGGGA'	TAACES M.avium
2924	GGTGTCACTCAACGG	ATAAAAGGT.	ACCCCGGGGA'	TAACAG M.paratuberc.
3625	GGTGTCGCTCAACGG	ATAAAAGGT:	ACCCCGGGGA'	TAACAG M.tuberculosis
3017	GGTGTCGCTCAACGG	ATAAAAGGT:	ACCCCGGGGA'	FAACAG M.phlei
2948	GGTGTCGCTCAACGG	ATAAAAGGT	accccggga!	TAACAG M.leprae
1910				M.gastri
2652	GGTGTCGCTCAACGG	ATAAAAGGT	accccggga!	TAACAG M.kansasii
5102	GGTGTCGCTCAACGG	ATAAAAGGT	ACCCCGGGGA!	TAACAG M.smegmatis
				-
	2730	2740	2750	2760
2064				
2964	GCTGATCTTCCCCAA	GAGTCCATA'	rcgacgggat(GGTTTG M.avium
2504	GCTGATCTTCCCCAA	GAGTCCATA'	rcgacgggat(GGTTTG M.paratuberc.
3003	GOTGATOTTCCCCAA	GAGTCCATA!	rcgacgggat(GGTTTG M.tuberculosis
305/	GCTGATCTTCCCCAA	GAGTCCATA'	rcgacgggato	GGTTTG M.phlei
1010	GCTGATCTTCCCCAA	GAGTCCATA!	rcgacgggate	
1910	CCMC b mcmmccccc >	03.0mc		M.gastri
209Z	CCTGATCTTCCCCAA	GAGTCCATA!	rcgacgggate	GGTTTG M.kansasii
3142	GUTGATUTTUCCCCAA	GAGTCCATA!	ICGACGGGATO	GTTTG M.smegmatis

Figure 4K

					
	27	70	2780	2790	2800
3004	GCACCTCGA!	TGTCGGCT	CGTCGCATC	CTGGGGCTGG	AGCA M.avium
3004	GCACCTCGA!	rgtcggci	CGTCGCATC	CTGGGGCTGG	AGCA M.paratuberc.
3705	GCACCTCGA:	rgtcggct	'CGTCGCAT(CTGGGGCTGG	AGCA M.tuberculosis
3097	GCACCTCGA:	rgtcggci	CGTCGCATC	CTGGGGCTGG	AGCA M.phlei
3028	GCACCTCGA!	rgtcggci	CGTCGCATC	CTGGGGCTGA	AGCA M.leprae
1910				_	M_dastri
2732	GCACCTCGA!	rgtcggct	CGTCGCATC	CTGGGGCTGG	AGCA M.kansasii
5182	GCACCTCGAT	rgtcggct	CGTCGCATC	CTGGGGCTGG	AGCA M.smegmatis
	28	10	2820	2830	2840
3044	GGTCCCNIN	CONTRACTOR	TCTTCCCCC	1 DEMPERATOR	GCAC M.avium
3044	GGTCCCATEG	1000C	TGTTCGCCC	-ATTAAAGCGG	GCAC M.avium GCAC M.paratuberc.
3745	GGTCCCAAGG	76911666C	TGTTCGCCC	-ATTAAAGCGG	GCAC M.paratuberc. GCAC M.tuberculosis
3137	GGTCCCAAGG	CTTGGGC	TGTTCGCCC	-ATTAAAGCGG	GCAC M.tuberculosis GCAC M.phlei
3088	GGTCCCAAGG		TGTTCGCCC	-ATTAAAGCG	GCAC M.phlei GCAC M.leprae
1910	ag I C C C ANDIG	99119990	IGITCGCCC	-ATTAAAGCGG	
	GGTCCCNNGG	יכתייככככ	пстисссо		M.gastri GCAC M.kansasii
5222	GGTCCCAAGG	:CTTCCCC	TGTTCGCCC		GCAC M.Kansasıı GCAC M.smegmatis
3222	GGICCCAME	99119990	refredece	UATTAAAGCG0	CAC M.smegmatis
	•				
	•				
	305		3060	3070	2000
2202	305	· -	3060	· -	3080
3283	CAAGATCAGG	TTT-CTC	CCCTTTTAG	AEGGATAAGGC	CCC M. avium
638	CAAGATCAGG	TTT-CTCA	CCCTTTTAG	AGGGATAAGGC	CCC M.avium
638 3283	CAAGATCAGG CAAGATCAGG CAAGATCAGG	TTT-CTCA TTT-CTCA	CCCTTTTAG CCCTTTTAG	AGGGATAAGGC	CCC M.avium
638 3283 3984	CAAGATCAGG CAAGATCAGG CAAGATCAGG CAAGATCAGG	TTT-CTCF TTT-CTCF TTT-CTCF TTT-CTCF	CCCTTTTAG CCCTTTTAG CCCTTTTAG CCCAGTTG	ABGGATAAGGC AGGGATAAGGC AGGGATAAGGC AGGGATAAGGC	CCC M.avium CCC M.intracellulare CCC M.paratuberc.
638 3283 3984 570	CAAGATCAGG CAAGATCAGG CAAGATCAGG CAAGATCAGG CAAGATCAGG	TTT-CTCA TTT-CTCA TTT-CTCA TTT-CTCA TTT-CTCA	CCCTTTTAG CCCTTTTAG CCCTTTTAG CCCACTTGG	EGGATARGO ARGESTARGO A	CCC M.avium CCC M.intracellulare CCC M.paratuberc. CCC M.tuberculosis
638 3283 3984 570 3376	CAAGATCAGG CAAGATCAGG CAAGATCAGG CAAGATCAGG CAAGATCAGG CAAGATCAGG	TTT-CTCA TTT-CTCA TTT-CTCA TTT-CTCA TTT-CTCA	CCCTTTTAG CCCTTTTAG CCCTTTTAG CCCACTTGG	ABGGATAAGGC AGGGATAAGGC AGGGATAAGGC AGGGATAAGGC	CCC M.avium CCC M.intracellulare CCC M.paratuberc. CCC M.tuberculosis CCC M.bovis CCC M.phlei
638 3283 3984 570 3376 3307	CAAGATCAGG CAAGATCAGG CAAGATCAGG CAAGATCAGG CAAGATCAGG CAAGATCAGG	TTT-CTCA TTT-CTCA TTT-CTCA TTT-CTCA TTT-CTCA	CCCTTTTAG CCCTTTTAG CCCTTTTAG CCCACTTGG	EGGATARGO ARGESTARGO A	CCC M.avium CCC M.intracellulare CCC M.paratuberc. CCC M.tuberculosis CCC M.bovis CCC M.phlei M.leprae
638 3283 3984 570 3376 3307 1910	CAAGATCAGG CAAGATCAGG CAAGATCAGG CAAGATCAGG CAAGATCAGG CAAGATCAGG CAAGATCAGG CAAGATCAGG	TTT-CTCA TTT-CTCA TTT-CTCA TTT-CTCA TTT-CTCA	ACCUTTTACO ACCUTTTACO ACCUTTTACO ACCUTTTACO ACCUTTCACO ACCUTT	EGGATANGGO SAGGGATANGGO SAGGATANGGO TGGGATANGGO TGGGATANGGO SAGGGATANGGO	CCC M.avium CCC M.intracellulare CCC M.paratuberc. CCC M.tuberculosis CCC M.bovis CCC M.phlei M.leprae
638 3283 3984 570 3376 3307 1910 3011	CAAGATCAGG CAAGATCAGG CAAGATCAGG CAAGATCAGG CAAGATCAGG CAAGATCAGG	TTT-CTCA TTT-CTCA TTT-CTCA TTT-CTCA GTT-CTCA	CCQTTTTAG CCTTTTAG CCACTTGG CCACTGG CCACTGGG CCACTGTCGG	EGGATAAGGO SAGGGATAAGGO SAGGATAAGGO TGGGATAAGGO SAGGGATAAGGO	CCC M.avium CCC M.intracellulare CCC M.paratuberc. CCC M.tuberculosis CCC M.bovis CCC M.phlei M.leprae M.gastri
638 3283 3984 570 3376 3307 1910 3011	CAAGATCAGG CAAGATCAGG CAAGATCAGG CAAGATCAGG CAAGATCAGG CAAGATCAGG	TTT-CTCA TTT-CTCA TTT-CTCA TTT-CTCA GTT-CTCA	CCQTTTTAG CCTTTTAG CCACTTGG CCACTGG CCACTGGG CCACTGTCGG	EGGATAAGGO SAGGGATAAGGO SAGGATAAGGO TGGGATAAGGO SAGGGATAAGGO	CCC M.avium CCC M.intracellulare CCC M.paratuberc. CCC M.tuberculosis CCC M.bovis CCC M.phlei M.leprae
638 3283 3984 570 3376 3307 1910 3011	CAAGATCAGG CAAGATCAGG CAAGATCAGG CAAGATCAGG CAAGATCAGG CAAGATCAGG	TTT-CTCA TTT-CTCA TTT-CTCA TTT-CTCA GTT-CTCA	CCQTTTTAG CCTTTTAG CCACTTGG CCACTGG CCACTGGG CCACTGTCGG	EGGATAAGGO SAGGGATAAGGO SAGGATAAGGO TGGGATAAGGO SAGGGATAAGGO	CCC M.avium CCC M.intracellulare CCC M.paratuberc. CCC M.tuberculosis CCC M.bovis CCC M.phlei M.leprae M.gastri
638 3283 3984 570 3376 3307 1910 3011	CAAGATCAGG CAAGATCAGG CAAGATCAGG CAAGATCAGG CAAGATCAGG CAAGATCAGG	TTT-CTCA TTT-CTCA TTT-CTCA TTT-CTCA GTT-CTCA	CCQTTTTAG CCTTTTAG CCACTTGG CCACTGG CCACTGGG CCACTGTCGG	EGGATAAGG SAGGGATAAGG SAGGGATAAGG TGGGATAAGG TGGGATAAGG SAGGGATAAGG TGGGATAAGG SAGGGATAAGG	CCC M.avium CCC M.intracellulare CCC M.paratuberc. CCC M.tuberculosis CCC M.bovis CCC M.phlei M.leprae M.gastri
638 3283 3984 570 3376 3307 1910 3011 5462	CAAGATCAGG CAAGATCAGG CAAGATCAGG CAAGATCAGG CAAGATCAGG CAAGATCAGG CAA	TTT-CTCA TTT-CTCA TTT-CTCA TTT-CTCA GTT-CTCA GTT-CTCA	CCCTTTTAG CCCTTTTAG CCCACTTGG CCCACTTGG CCCACTTAGG CCCACTTAGG	EGGATAAGGC FAGGGATAAGGC FAGGGATAAGGC FAGGGATAAGGC FAGGGATAAGGC FAGGGATAAGGC FAGGGATAAGGC FAGGGATAAGGC	CCC M.avium CCC M.intracellulare CCC M.paratuberc. CCC M.tuberculosis CCC M.bovis CCC M.phlei M.leprae M.gastri CCC M.kansasii CCC M.smegmatis
638 3283 3984 570 3376 3307 1910 3011 5462	CAAGATCAGG CAAGATCAGG CAAGATCAGG CAAGATCAGG CAAGATCAGG CAAGATCAGG CAAGATCAGG CAAGATCAGG CAAGATCAGG	TTT-CTCA TTT-CTCA TTT-CTCA TTT-CTCA GTT-CTCA GTT-CTCA	CCCTTTTAG CCCTTTTAG CCCACTTGG CCCACTTGG CCCACTTGG CCCACTTGGG CCCACTTGGG	EAGGATAAGGCEAGGGATAAGGCEAGGATAAGGCEATAAGGCEATAAGGCEATAAGGCEATAAGGCEAGGATAAGGCEAGGGATAAGGCEAGGGATAAGGCEAGGGATAAGGCEAGGGATAAGGCEAGGGATAAGGCEAGGGATAAGGCEAGGGATAAGGCEAGGGATAAGGCEAGGGATAAGGCEAGGGATAAGGCEAGGGATAAGGCEAGGGATAAGGCEAGGGATAAGGCEAGGGATAAGGCEAGGGATAAGGCEAGGGATAAGGCEAGGATAAGGCEAGGGATAAGGCEAGGGATAAGGCEAGGGATAAGGCEAGGGATAAGGCEAGGAGGATAAGGCEAGGAAGGCATGGAAGGCATGGAAGGACCTGGAAGGAAGGCATAAGGCAAGGCATGGAAGGACCTGGAAGGAA	CCC M.avium CCC M.intracellulare CCC M.paratuberc. CCC M.tuberculosis CCC M.bovis CCC M.phlei M.leprae M.gastri CCC M.kansasii CCC M.smegmatis CCC M.smegmatis
638 3283 3984 570 3376 3307 1910 3011 5462	CAAGATCAGG CAAGATCAGG CAAGATCAGG CAAGATCAGG CAAGATCAGG CAAGATCAGG CAAGATCAGG CAAGATCAGG CAAGATCAGG CCCGC-AGACCCCCGC-AGACCCCCGC-AGACCCCCCCCC	TTT-CTCA TTT-CTCA TTT-CTCA TTT-CTCA CTT-CTCA CTT-CTCA CTT-CTCA CTCACCACCACCACCACCACCACCACCACCACCACCACCA	CCCTTTTAG CCCTTTTAG CCCACTTGG CCCACTTGG CCCACTTGG CCCACTTGGG CCCACTTGGG CCCTCTAGG	EAGGATAAGGCEAGGATAAGGCEAGGATAAGGCEATAAGGCEATAAGGCEATAAGGCEATAAGGCEAGGATAAGGCEAGGATAAGGCEAGGATAAGGCEAGGATAAGGCEAGGATAAGGCEAGGACCTGGAAGGACCTGGAAGGACCTGGAAGGACCTGGAAGGACCTGGAAGGACCTGGAAGGACCTGGAAGGACCTGGAAGGACCTGGAAGGACCTGGAAG	CCC M.avium CCC M.intracellulare CCC M.paratuberc. CCC M.tuberculosis CCC M.bovis CCC M.phlei M.leprae M.gastri CCC M.kansasii CCC M.smegmatis CCC M.avium CCT M.avium CCT M.intracellulare
638 3283 3984 570 3376 3307 1910 3011 5462 3322 677 3322	CAAGATCAGG CAAGATCAGG CAAGATCAGG CAAGATCAGG CAAGATCAGG CAAGATCAGG CAAGATCAGG CAAGATCAGG CAAGATCAGG CCAAGATCAGG CCCGC-AGACCCCGC-AGACCCCGC-AGACCCCCCCCCC	TTT-CTCA TTT-CTCA TTT-CTCA TTT-CTCA CTT-CTCA CTT-CTCA CTCACCACACACACACACACACACACACACACACA	CCCTTTTAG CCCTTTTAG CCCACTTGG CCCACTTGG CCCACTTGG CCCACTTGGG CCCACTTAGG CCCTCTAGG TGATAGGCC	EAGGEATAAGGCEAGGGATAAGGCEAGGGATAAGGCEATAAGGCEATAAGGCEATAAGGCEAGGGATAAGGCEAGGGATAAGGCEAGGGATAAGGCEAGGCATGGAAGGCCTGGAAGGACCTGGAAGGACCTGGAAGGACCTGGAAGGACCTGGAAG	CCC M.avium CCC M.intracellulare CCC M.paratuberc. CCC M.tuberculosis CCC M.bovis CCC M.phlei M.leprae M.gastri CCC M.kansasii CCC M.smegmatis CCC M.avium CCT M.avium CCT M.avium CCT M.paratuberc
638 3283 3984 570 3376 3307 1910 3011 5462 3322 677 3322 4023	CAAGATCAGG CAAGATCAGG CAAGATCAGG CAAGATCAGG CAAGATCAGG CAAGATCAGG CAAGATCAGG CAAGATCAGG CAAGATCAGG CCAAGATCAGG CCAAGATCAGG CCAGCAGACCCCGCAGACCCCGCAGACCCCCGCAGACCCCCC	TTT-CTCA TTT-CTCA TTT-CTCA TTT-CTCA CTT-CTCA CTT-CTCA CTCACGGGTT CACGGGTT CACGGGTT CACGGGTT	CCCTTTTAG CCCTTTTAG CCCACTTGG CCCACTTGG CCCACTTGG CCCTCTAGG CCCTCTAGG CCCTCTAGG CCCTCTAGG CCCTCTAGG CCCTCTAGG CCCTCTAGGC CAATAGGCC	EGGATAAGGC EAGGGATAAGGC EAGGGATAAGGC EAGGGATAAGGC EAGGGATAAGGC EAGGGATAAGGC EAGGGATAAGGC EAGGGATAAGGC EAGGGATAAGGC EAGGGATAAGGC EAGACCTGGAAG	CCC M.avium CCC M.intracellulare CCC M.paratuberc. CCC M.tuberculosis CCC M.bovis CCC M.phlei M.leprae M.gastri CCC M.kansasii CCC M.smegmatis CCC M.avium CCT M.avium CCT M.intracellulare CCT M.paratuberc. CCT M.tuberculosis
638 3283 3984 570 3376 3307 1910 3011 5462 3322 677 3322 4023 609	CAAGATCAGG CAAGATCAGG CAAGATCAGG CAAGATCAGG CAAGATCAGG CAAGATCAGG CAAGATCAGG CAAGATCAGG CAAGATCAGG CCAAGATCAGG CCAGCAGACCCCGCAGACCCCGCAGACCCCCGCAGACCCCCC	TTT-CTCA TTT-CTCA TTT-CTCA TTT-CTCA CTT-CTCA CTT-CTCA CACGGGTT CACGGGTT CACGGGTT CACGGGTT CACGGGTT	CCCTTTTAG CCCTTTTAG CCCACTTGG CCCACTTGG CCCACTTGG CCCACTTGG CCCTCTAGG CCCTCTAGG CCCTCTAGG CCCTCTAGG CCATAGGCC CAATAGGCC CAATAGGTC	EGGATAAGGC EAGGGATAAGGC EAGGGATAAGGC EAGGGATAAGGC EAGGGATAAGGC EAGGGATAAGGC EAGGGATAAGGC EAGGGATAAGGC EAGGGATAAGGC EAGGGATAAGGC EAGACCTGGAAGC EAGACCTGGAAGC	CCC M.avium CCC M.intracellulare CCC M.paratuberc. CCC M.tuberculosis CCC M.bovis CCC M.phlei M.leprae M.gastri CCC M.kansasii CCC M.smegmatis CCC M.avium CCT M.avium CCT M.intracellulare CCT M.paratuberc. CCT M.tuberculosis CCT M.bovis
638 3283 3984 570 3376 3307 1910 3011 5462 3322 677 3322 4023 609	CAAGATCAGG CAAGATCAGG CAAGATCAGG CAAGATCAGG CAAGATCAGG CAAGATCAGG CAAGATCAGG CAAGATCAGG CAAGATCAGG CCAAGATCAGG CCAGCAGACCCCGCAGACCCCGCAGACCCCCGCAGACCCCCC	TTT-CTCA TTT-CTCA TTT-CTCA TTT-CTCA CTT-CTCA CTT-CTCA CACGGGTT CACGGGTT CACGGGTT CACGGGTT CACGGGTT	CCCTTTTAG CCCTTTTAG CCCACTTGG CCCACTTGG CCCACTTGG CCCACTTGG CCCTCTAGG CCCTCTAGG CCCTCTAGG CCCTCTAGG CCATAGGCC CAATAGGCC CAATAGGTC	EGGATAAGGC EAGGGATAAGGC EAGGGATAAGGC EAGGGATAAGGC EAGGGATAAGGC EAGGGATAAGGC EAGGGATAAGGC EAGGGATAAGGC EAGGGATAAGGC EAGGGATAAGGC EAGACCTGGAAG	M. avium CCC M.intracellulare CCC M.paratuberc. CCC M.tuberculosis CCC M.bovis CCC M.phlei M.leprae M.gastri CCC M.kansasii CCC M.smegmatis CCC M.avium CCT M.avium CCT M.intracellulare CCT M.paratuberc. CCT M.tuberculosis CCM M.bovis CCM M.phlei
638 3283 3984 570 3376 3307 1910 3011 5462 3322 677 3322 4023 609 3415	CAAGATCAGG CAAGATCAGG CAAGATCAGG CAAGATCAGG CAAGATCAGG CAAGATCAGG CAAGATCAGG CAAGATCAGG CAAGATCAGG CCAAGATCAGG CCAGCAGACCCCGCAGACCCCGCAGACCCCCGCAGACCCCCC	TTT-CTCA TTT-CTCA TTT-CTCA TTT-CTCA CTT-CTCA CTT-CTCA CACGGGTT CACGGGTT CACGGGTT CACGGGTT CACGGGTT	CCCTTTTAG CCCTTTTAG CCCACTTGG CCCACTTGG CCCACTTGG CCCACTTGG CCCTCTAGG CCCTCTAGG CCCTCTAGG CCCTCTAGG CCATAGGCC CAATAGGCC CAATAGGTC	EGGATAAGGC EAGGGATAAGGC EAGGGATAAGGC EAGGGATAAGGC EAGGGATAAGGC EAGGGATAAGGC EAGGGATAAGGC EAGGGATAAGGC EAGGGATAAGGC EAGGGATAAGGC EAGACCTGGAAGC EAGACCTGGAAGC	M. avium CCC M.intracellulare CCC M.paratuberc. CCC M.tuberculosis CCC M.bovis CCC M.phlei M.leprae M.gastri CCC M.kansasii CCC M.smegmatis CCC M.avium CCT M.avium CCT M.intracellulare CCT M.paratuberc. CCT M.tuberculosis CCT M.bovis CCT M.bovis CCT M.phlei M.leprae
3322 677 3322 4023 609 3415 3309 1910 3050	CAAGATCAGG CAAGATCAGG CAAGATCAGG CAAGATCAGG CAAGATCAGG CAAGATCAGG CAAGATCAGG CAAGATCAGG CAAGATCAGG CCAGCAGACCAGAC	TTT-CTCA TTT-CTCA TTT-CTCA TTT-CTCA TTT-CTCA TTT-CTCA CTT-CTCA CTCACGGGTT CACGGGTT CACGGTT CAC	CCCTTTTAG CCCTTTTAG CCCACTTGG CCCACTTGG CCCACTTGG CCCACTTGG CCCACTGGGC CCCACTGGGC CCCACTGGGCC CACTGGGCC CACTGGGCC CACTGGGCC CACTGGGCC CACTGGGCC CACTGGGCC	EAGACCTGGAAGGCCAGACCTGGAAGGCCTGGAAGGCCTGGAAGGCCTGGAAGGCCTGGAAGGCCTGGAAGCCTGAAGCAAGC	CCC M.avium CCC M.intracellulare CCC M.paratuberc. CCC M.tuberculosis CCC M.bovis CCC M.phlei M.leprae M.gastri CCC M.kansasii CCC M.smegmatis CCT M.avium CCT M.intracellulare CCT M.paratuberc. CCT M.tuberculosis CCT M.bovis CCT M.bovis CCT M.phlei M.leprae M.gastri M.leprae M.gastri CCT M.kansasii
3322 677 3322 4023 609 3415 3309 1910 3050	CAAGATCAGG CAAGATCAGG CAAGATCAGG CAAGATCAGG CAAGATCAGG CAAGATCAGG CAAGATCAGG CAAGATCAGG CAAGATCAGG CCAGCAGACCAGAC	TTT-CTCA TTT-CTCA TTT-CTCA TTT-CTCA TTT-CTCA TTT-CTCA CTT-CTCA CTCACGGGTT CACGGGTT CACGGTT CAC	CCCTTTTAG CCCTTTTAG CCCACTTGG CCCACTTGG CCCACTTGG CCCACTTGG CCCACTGGGC CCCACTGGGC CCCACTGGGCC CACTGGGCC CACTGGGCC CACTGGGCC CACTGGGCC CACTGGGCC CACTGGGCC	EAGACCTGGAAGGCCAGACCTGGAAGGCCTGGAAGGCCTGGAAGGCCTGGAAGGCCTGGAAGGCCTGGAAGCCTGAAGCAAGC	M. avium CCC M.intracellulare CCC M.paratuberc. CCC M.tuberculosis CCC M.bovis CCC M.phlei M.leprae M.gastri CCC M.kansasii CCC M.smegmatis CCC M.avium CCT M.avium CCT M.intracellulare CCT M.paratuberc. CCT M.tuberculosis CCT M.bovis CCT M.bovis CCT M.phlei M.leprae

Figure 4L

		130	140	150	16	O .
107	GAGTAACAG	стесств	יייבייפר כריי	GCACTTC-GG	ממתמט	M. cardam
59	GAGTAACAC	CGTGGGCA	ATCTGCCCT	CCACTIC GG	CULUVY	M.intracellulare
107	GAGTAACAG	CGTGGGCAI	11010001 110100001	GCACTTC GG	תמשתם! מאדאא	M. naratuberc.
70	GAGTAACAG	CTGGGCA	,TCTECCCT	COVOLIC GG	CULLY	M.scrofulaceum
70	GAGTAACAG	corecercia	TCTGCCCT	GCACTIC GG	CATAR	M.tuberculosis
209	GAGTAACAC	COTGGGTG	\TCTGCCCT	GCACTTC-GG	CATAA	M bouis
120	GAGTAACAC	CTGCCTA	TCTGCCCT	GCACTTCAGG	CATAA	M lennes
69	GAGTAACAC	CGTGGGCAR	.TCTGCCCT	CACACACC	CATAA	M.kansasii
70	GAGTAACAC	CGTGGGCAR	TCTGCCCT	GCACACC-GG	CATAA	M. castri
104	GAGTAACAC	ствевпая	TCTGCCCT	CACATTC-CC	CATAR	M.gordonae
64	GAGTAACAC	стессси	TCTGCCCT	607 077 10 66	CATAN	M.marinum
		, c 1 c c c c c c		JOACIIC GG	GNIAM	M.Marinum
	•					
		F 0	150	420	1	
		50	460	470	48	_
424	AAACCTCTI	TCACCATO	GACGAAGG	TCCGGC TTTT	CTCGG	M.avium
376	AAACCTCTT	TCACCATC	GACGAAGG	TCCGGGTTTT	CTCGG	M.intracellulare
424	AAACCTCTT	TCACCATC	GACGAAGG	TCCGGGTTTT	CTAGG	M.paratuberc.
387	AAACCTCTT	TCACCATC	GACGAAGG	CTCA drr	TGTGG	M.scrofulaceum
389	AAACCTCTT	'TCACCATC	GACGAAGG	TCCGGGTTCT	CTCGG	M.tuberculosis
528	AAACCTCTT	TCACCATC	GACGAAGG	TCCGGGTTCT	CTCGG	M howin
439	AAACCTCTT	TCACCATC	GACGAAGG	TCTGGGAATT	CTCGG	M.leprae
386	AAACCTCTT	TCACCATC	GACGAAGG'	TCCGGGTTCT	CTCGG	M.kansasii
387	AAACCTCTT	TCACCATC	GACGAAGG	TCCGGGTTdT	CTCGG	M.gastri
420	AAACCTCTT	TCACCATC	GACGAAGG	TCCGGGTTTT	CTCGG	M.gordonae
381	AAACCTCTT	TCACCATC	GACGAAGG	TICGGGTTTT	CTCGG	M.marinum
		•		_		
					· · ·	
	49		500	510	520	
129	ATTGACGGTA	GGTGGAGA	AGAAGCAC	CCGCCAACT	ACGTG	M.tuberculosis
568	ATTGACGGTA	GGTGGAG	AGAAGCA(CCGCCCAACT	ACGTG	M.bovis
164	ATTGACGGTA	lggtggag <i>f</i>	AGAAGCAC	CCGCCAACT	ACGTG	M.avium
416	ATTGACGGTA	GGTGGAGA	AGAAGCAC	CCGCCAACT	ACGTG	M.intracellulare
164	ATTGACGGTA	GGTGGAGA	AGAAGCAC	ACT	ACGTG	M.paratuberc.
124	GTTGACGGTA	GGTGGAG	AGAAGCAC	CGGCCAACT	ACGTG	M.scrofulaceum
179	ATTGACGGTA	GGTGGAGF	AGAAGCAC	CGGCCAACT	ACGTG	M.leprae
126	ATTGACGGTA	GGTGGAGF	AGAAGCAC	CGGCCAACT	ACGTG	M.kansasii
127	ATTGACGGTA	GGTGGAGA	AGAAGCAC	CGGCCAACT	ACGTG	M.gastri
160	GOTGACGGTA	GGTGGAGF	AGAAGCAC	CCGCCAACT	ACGTG	M.gordonae
						yor donae

Figure 5A

	1130	1140	1150	1160
1104	TCTCATGTTGCCA	GOGGTAATGC	GGGGACTCG'	TGAGAG M.avium
1056	TCTCATGTTGCCA	GCGGGTAATGCC	GGGGACTCG'	TGAGAG M.intracellulare
1098	TCTCATGTTGCCA	GCGGGTAATGCA	GGGGACTCG	TGAGAG M.paratuberc.
1064	TCTCATGTTGCCA	GCGGGTAATGCC	GGGGACTCG'	TGAGAG M.scrofulaceum
1069	TCTCATGTTGCCA	GCACGTAATGGT	GGGGACTCG'	TGAGAG M.tuberculosis
1208	TCTCATGTTGCCA	SCACGTAATGGT	GGGGACTCG'	TGAGAG M hovis
1119	TCTCATGTTGCCA	GCACGTAATGGT	GGGGACTCG'	TGAGAG M.leprae
1066	TCTCATGTTGCCA	SCGGGTAATGCC	GGGGACTCG'	rgagag M.kansasii
1067	TCTCATGTTGCCA	SCGGGTAATGCC	GGGGACTCG'	rgagag M.gastri
1100	TCTCATGTTGCCA	GCGGGTAATGCC	GGGGACTCG'	rgagag M.gordonse
1061	TCTCATGTTGCCA	SCACGTAATGGT	GGGGACTCG	rgagag M.marinum
		·		
	1290	1300	1310	1320
1264	CGAATCCTTTTAA	AGCCGGACTCAG	TTCGGATTGG	GGTCT M. avium
1216	CGAATCCTTTTAA	AGCCGGTCTCAG	TTCGGATTGG	GGTCT M.intracellulare
1258	CGAATCCTTTTAA	AGCCGGACTCAG	TTCGGATTGG	GGGTCT M.paratuberc.
1224	CGAATCCTTTTAAA	AGCCGGTCTCAG	ттсаватСве	GGTCT M scrofulaceum
1229	CGAATCCTTA-AAA	AGCCGGTCTCAG	TTCGGATCG	GGGTCT M.tuberculosis
1368	CGAATCCTTA-AAJ	AGCCGGTCTCAG	TTCGGATCG	GGTCT M.bovis
1279	CGAATCCTTTTAA	AGCCGGTCTCAG	TTCGGATCG	GGTCT M.leprae
1226	CGAATCCTTTTAA	AGCCGGTCTCAG'	TTCGGATCG	GGGTCT M.leprae GGGTCT M.kansasii
1227	CGAATCCTTTTAAI	AGCCGGTCTCAG'	PTCGGATICG	GGTCT M destri
1260	CGAATCCTTTTAA	agccgg ctcag	rtcggatcg	GGTCT M.gordonae
1221	CGAATCCTTT	agccgg[[ctcag:	TTCGGATCGG	GGGTCT M.marinum
				
	1330	1340	1350	1360
	GCAACTCGACCCC			
1256	GCAACTCGACCCCA	TGAAGTCGGAG	rcgctagta <i>p</i>	TCGCA M.intracellulare
1298	GCAACTAGACCOA	TGAAGTCGGAG'	rcgctagta <i>p</i>	ATCGCA M.paratuberc.
1264	GCAACTCGACCCC	TGAAGTCGGAG'	rcgctagtaf	TCGCA M.scrofulaceum
1268	GCAACTCGACCCC	TGAAGTCGGAG:	CGCTAGTAP	TCGCA M.tuberculosis
1407	GCAACTCGACCCC	TGAAGTCGGAG'	CGCTAGTAP	ATCGCA M.bovis
1319	GCAACTCGACCCC	TGAAGTCGGAG:	CGCTAGTAP	TCGCA M.leprae
1266	GCAACTCGACCCC	TGAAGTCGGAG'	rcgctagta _l a	TCGCA M.kansasii
1267	GCAACTCGACCCC	TGAAGTCGGAG'	rcgctagta <i>p</i>	TCGCA M.gastri
1300	GCAACTCGACCCC	TGAAGTCGGAG:	rcgctagtaa	TCGCA M.gordonae
1260	GCAACTCGACCCC	TGAAGTCGGAG'	CGCTAGTAP	TCGCA M.marinum

Figure 5B

Figure 6

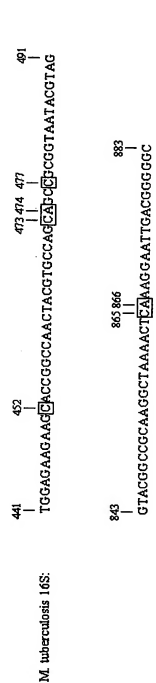


Figure 7

INTERNATIONAL SEARCH REPORT

Inter anal Application No PCT/DK 97/00425

		FC	1/08 9//00425	
A. CLASSI IPC 6	IFICATION OF SUBJECT MATTER C12Q1/68 C07K14/00			
According to	o international Patent Classification (IPC) or to both national classific	ation and IPC		
	SEARCHED			
Minimum do IPC 6	ocumentation searched (classification system followed by classification C12Q C07K	on symbols)		
Documenta	tion searched other than minimum documentation to the extent that s	uch documents are included in	the fields searched	
Electronic d	lata base consulted during the international search (name of data ba	se and, where practical, searc	h terma used)	
C. DOCUM	ENTS CONSIDERED TO BE RELEVANT			
Category •	Citation of document, with indication, where appropriate, of the rele	evant passages	Relevant to claim No.	
Υ	US 5 547 842 A (HOGAN JAMES ET AL) 20 August 1996 cited in the application see the whole document		1-36	
Ý	WO 96 17956 A (GENE POOL INC ;WEININGER SUSAN (US); WEININGER ARTHUR M (US)) 13 June 1996 see the whole document		1-36	
Υ .	WO 95 32305 A (DAKO AS) 30 November 1995 see the whole document		1-36	
Α	EP 0 572 120 A (GEN PROBE INC) 1 December 1993 cited in the application see the whole document			
		-/		
Y Further documents are listed in the continuation of box C.		X Patent family member	rs are listed in annex.	
Special categories of cited documents: "A" document defining the general state of the art which is not		or priority date and not in	after the international filing date conflict with the application but	
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Date of the a	actual completion of theinternational search	Date of mailing of the international search report		
20 January 1998		30/01/1998		
Name and mailing address of the ISA European Patent Office, P.B. 5818 Patentlaan 2 NL - 2280 HV Rijswijk		Authorized officer		
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212		PCT/DK 97/00425
C.(Continu	Atlon) DOCUMENTS CONSIDERED TO BE RELEVANT Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
	- in the second	risisvali to Gauti No.
P,Y	WO 96 36734 A (ABBOTT LAB) 21 November 1996 see the whole document	1-36
		<u>.</u>
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INTERNATIONAL SEARCH REPORT

Information on patent family members

Inter. July Application No PCT/DK 97/00425

Patent document cited in search report	Publication date	Patent family member(s)	Publication date
US 5547842 A	20-08-96	US 5541308 A	30-07-96
		US 5595874 A	
		US 5593841 A	
		US 5683876 A	
		US 5677127 A	
		US 5677128 A	
		US 5677129 A	
		US 5693468 A	
		US. 5691149 A	
		US 5693469 A	
		US 5679520 A	
		US 5674684 A	
		DK 413788 A	
		AU 616646 B	
		AU 1041988 A	
•		EP 0272009 A	
		JP 1503356 T	
		KR 9511719 B	
		WO 8803957 A	
WO 9617956 A	13-06-96	AU 4418996 A	26-06-96
		CA 2206127 A	
		EP 0796344 A	24-09-97
		NO 972611 A	11-08-97
WO 9532305 A	30-11-95	AU 2522095 A	18-12-95
,		EP 0760008 A	05-03-97
EP 0572120 A	01-12-93	AU 4114793 A	29-11-93
		JP 7506723 T	
		WO 9322330 A	
WO 9636734 A	21-11-96	NONE	——————————————————————————————————————

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